NOTE to Phyllis: Apparently "Acivacin" was misspelled in the claims. It's not Registry, and CAPlus contains only one record for a compound that may be the same, and it's spelled "Acivicin".

=> d hit acivacin all

1-6 (Pharmacology)

L22 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2005 ACS on STN 1983:27436 HCAPLUS AN 98:27436 DN Entered STN: 12 May 1984 ED Biochemical pharmacology of acivicin in rat hepatoma cells TILui, May S.; Kizaki, Harutoshi; Weber, George ΑU Sch. Med., Indiana Univ., Indianapolis, IN, 46223, USA Biochemical Pharmacology (1982), 31(21), 3469-73 CS SO CODEN: BCPCA6; ISSN: 0006-2952 DTJournal English LΑ

CC

GI

AB The antiglutamine agent acivicin (I) [42228-92-2] inhibited the growth of hepatoma 3924A cells in culture. The 7 day LC50 of acivicin was determined to be 1.4 µM. A combination of cytidine, deoxycytidine, and guanosine completely protected the hepatoma cells against the cytotoxicity acivicin, but each nucleoside by itself had no effect. Acivicin (0.1 mM) inhibited the incorporation of uridine and thymidine into macromols., but not that of leucine. Acivicin depressed the pools of CTP, GTP, dCTP, dGTP, and dTTP, but it increased the UTP levels. The activity of a highly purified CTP synthetase (EC 6.3.4.2) [9023-56-7] from rat liver and hepatoma 3924A was inhibited by acivicin. The inhibition was competitive with respect to L-glutamine, and the Ki values were 1.1 and 3.6 μM, resp. hydroxyacivicin [54549-02-9] Was also a competitive inhibitor, but it was less effective than acivicin, with a Ki value of 1.8 mM for the hepatoma enzyme. It appears that the principal mechanism of action of acivacin is the inhibition of CTP synthetase and GMP synthetase (EC 6.3.5.2) [37318-71-1]. acivicin antitumor biochem pharmacol ST Neoplasm inhibitors

IT

(hepatoma, acivicin, biochem. mechanism of)

IT 54549-02-9

RL: BIOL (Biological study)

(CTP synthetase inhibition by, hepatoma inhibition in relation to)

ΙT 42228-92-2

RL: BIOL (Biological study)

(hepatoma inhibition by, biochem. mechanism of)

ΙT 9023-56-7 37318-71-1

RL: BIOL (Biological study)

(inhibition of, by acivicin, hepatoma inhibition in relation to)

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L36 ANSWER 1 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER:
                     2005:248644 HCARLUS
DOCUMENT NUMBER:
                         142:274057
TITLE:
                         Sequences of human\schizophrenia related genes and use
                         for diagnosis, prognosis and therapy
                         Liew, Choong-chin
INVENTOR(S):
PATENT ASSIGNEE(S):
                                               Can.
                         Chondrogene Limited,
                         U.S. Pat. Appl. Publ., 156 pp., Cont.-in-part of U.S.
SOURCE:
                         Ser. No. 802,875.
                         CODEN: USXXCO
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                         KIND
                                             APPLICATION NO.
                                DATE
                                                                    DATE
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Searched by Mary Jane Ruhl

Ext. 22524

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PRIORITY APPLN. INFO.:
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                                                                     A2 20030620
                                                US 2004-802875
                                                                      A2 20040312
                                                US 2004 /812731
                                                                     A 20040330
     The present invention is directed to detection and measurement of gene
AΒ
```

The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring diseases using gene-specific and/or tissue-specific primers. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of 3 records for this document necessitated by the large number of index entries required to fully index the document and publication system constraints.].

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L36 ANSWER 2 OF 54 HCAPLUS COPYRIGHT 200 ACS on STN
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ACCESSION NUMBER: 2005:248643 HCAPLUS

DOCUMENT NUMBER: 142:274056

TITLE: Sequences of humann schizophrenia related genes and use

for diagnosis, prognosis and therapy

INVENTOR(S): Liew, Choong-Chi/n

PATENT ASSIGNEE(S): Chondrogene Limpted, Can.

SOURCE: U.S. Pat. Appl./ Publ., 156 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 2004241727	A1	2004 202	US 2004-812731		20040330 <
US 2004014059	A1	20040122	US 2002-268730		20021009 <
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US 2005196763	A1	200\$0908	US 2004-803857		20040318 <
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US 2004241727	A1	200/41202	US 2004-812731		20040330 <
PRIORITY APPLN. INFO.:			US 1999-115125P	P	19990106 <
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		- 1	US 2003-601518	A2	20030620
		1	US 2004-802875	A2	20040312
		1	US 2004-812731	Α	20040330

AB The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring diseases using gene-specific and/or tissue-specific

primers. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of 3 records for this document necessitated by the large number of index entries required to fully index the document and publication system constraints.].

L36 ANSWER 3 OF 54 HCAPLUS COPYRIGHT 2005 ACS of STN

ACCESSION NUMBER:

2005:172213 HCAPLUS

DOCUMENT NUMBER:

142:259426

TITLE:

Gene expression profiles and biomarkers for the

detection of asthma-replated and other disease-related

gene transcripts in blood

INVENTOR(S):

Liew, Choong-Chin

PATENT ASSIGNEE(S):

Chondrogene Limited, Can.

SOURCE:

U.S. Pat. Appl. Pub., 156 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 2005042630	A1	20050224	US 2004-816357		20040401 <
US 2004014059	A1	20040122	US 2002-268730		20021009 <
US 2005191637	A1	20050901	US 2004-803737		20040318 <
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PRIORITY APPLN. INFO.:		- 1	US 1999-115125P	P	19990106 <
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		- 1	US 2003-601518	A2	20030620
		1	US 2004-802875	A2	20040312
		- 1	US 2004-816357	A	20040401

The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing, and monitoring diseases, and in particular asthma, using gene-specific and/or tissue-specific primers. Affymetrix Human Genome U133 and ChondroChip microarrays were used to detect differentially expressed gene transcripts in hypertension, obesity, allergy, systemic steroids, coronary artery disease, diabetes type 2, hyperlipidemia, lung disease, bladder cancer, rheumatoid arthritis, osteoarthritis, liver cancer

, schizophrenia, Chagas disease, asthma, and manic depression syndrome. The present invention describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of three records for this document necessitated by the large number of index entries required to fully index the docoment and publication system constraints.].

L36 ANSWER 4 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 2005:160724 HCAPLUS

Searched by Mary Jane Ruhl Ext. 22524

DOCUMENT NUMBER: 142:259424

TITLE: Gene expression profiles and biomarkers for the

detection of asthma-related and other disease-related

gene transcripts in blood

INVENTOR(S): Liew, Choong-Chin

PATENT ASSIGNEE(S): Chondrogene Limited, Can.

SOURCE: U.S. Pat. Appl. Publ., 156 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE: LANGUAGE: Patent English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
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US 2005042630	A1	20050224	US 2004 / 816357		20040401 <
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			/US 2003-601518	A2	20030620
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			US 2004-816357	A	20040401

AB The present invention is directed to/detection and measurement of gene transcripts and their equivalent nugleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing, and monitoring diseases, and in particular asthma, using gene-specific and/or tissue-specific primers. Affymetrix Human Genome U133 and ChondroChip microarrays were used/to detect differentially expressed gene transcripts in hypertension, obestity, allergy, systemic steroids, coronary artery disease, diabetes type 2, hyperlipidemia, lung disease, bladder cancer, rheumatoid arthritis, osteoarthritis, liver cancer , schizophrenia, Chagas disease / asthma, and manic depression syndrome. The present invention describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of three records for this document necessitated by the large number of index enthies required to fully index the docoment and publication system constraints.].

L36 ANSWER 5 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:156228 HCAPLUS

Correction of: 2005:16967

DOCUMENT NUMBER: 142:192331

Correction of: 142:108390

TITLE: Quantitative RT-PCR method for the detection in blood

of microarray-identified rheumatoid arthritis-related gene transcripts for diagnosing and monitoring disease

state

INVENTOR(S): Liew, Choong-Chin

PATENT ASSIGNEE(S): Chondrogene Limited, Can.

SOURCE: U.S. Pat. Appl. Publ., 81 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
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US 2004265869	A1	20041230	US 2004- \$ 12716		20040330 <
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PRIORITY APPLN. INFO.:			US 199 ∮ -115125P	P	19990106 <
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			US 2903-601518		20030620
			US 2 / 04-802875	A2	20040312
			US 7 004-812782	Α	20040330

The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood for diagnosing and monitoring diseases. The present invention demonstrates that a simple drop of blood may be used to determine the quant. expression of various mRNAs that reflect the health/disease state of the subject through the use of quant. reverse transcription-polymerase thain reaction (QRT-PCR) anal. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring rheumatoid architis using gene-specific and/or tissue-specific primers. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen.

L36 ANSWER 6 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:155679 HCAPLUS

DOCUMENT NUMBER: 142:213366

TITLE: Quantitative RT-P¢R method for the detection in blood

of microarray-identified rheumatoid arthritis-related gene transcripts for diagnosing and monitoring disease

state

INVENTOR(S): Liew, Choong-Chir

PATENT ASSIGNEE(S): Chondrogene Limited, Can.

SOURCE: U.S. Pat. Appl. Publ., 81 pp., Cont.-in-part of U.S.

Ser. No. 802,875

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

Searched by Mary Jane Ruhl

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PRIORITY APPLN. INFO.:
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The present invention is directed to detection and measurement of gene AB transcripts and their equivalent nucleic acid products in blood for diagnosing and monitoring diseases. The present invention demonstrates that a simple drop of blood may be used to determine the quant. expression of various mRNAs that reflect the health/disease state of the subject through the use of quant. reverse transcription-polymerase chain reaction (QRT-PCR) anal. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring rheumatoid arthritis using gene-specific and/or tissue-specific primers. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease of to assess the effect of a particular treatment regimen. [This abstract record is one of 3 records for this document necessitated by the large/number of index entries required to fully index the document and publication system constraints.].

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L36 ANSWER 7 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN
                        2005:112850 HCAPLUS
ACCESSION NUMBER:
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DOCUMENT NUMBER:

142:153469

TITLE:

Gene expression profiles and biomarkers for the detection of Jung disease-related and other disease-related gene transcripts in blood

INVENTOR(S):

Liew, Choong-chin

PATENT ASSIGNEE(S):

Chondrogene Limited, Can.

SOURCE:

U.S. Pat. Appl. Publ., 155 pp., Cont.-in-part of U.S.

Ser. No. ,802,875. CODEN: UŚXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND/	DATE	APPLICATION NO.		DATE
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US 2004-812764 A 20040330

The present invention is directed to detection and measurement of gene AB transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring diseases using gene-specific and/or tissue-specific primers. Affymetrix Human Genome U133 and ChondroChip microarrays were used to detect differentially expressed gene transcripts in hypertension, obesity, allergy, systemic steroids, coronary artery disease, diabetes type 2, hyperlipidemia, lung disease, bladder cancer, rheumatoid arthritis, osteoarthritis, liver cancer, schizophrenia,/Chagas disease, asthma, and manic depression syndrome. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test/for disease or to assess the effect of a particular treatment regimen/ [This abstract record is one of 3 records for this document necessitated by the large number of index entries required to fully index the document/and publication system constraints.].

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L36 ANSWER 8 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STA
ACCESSION NUMBER: 2005:60759 HCAPLUS
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ACCESSION NUMBER: 2005:60759 HCAPLOS / Correction of: 2004:103657/2

DOCUMENT NUMBER: 142:111840

Correction of: 142:16824

TITLE: Gene expression profiles and biomarkers for the

detection of lung disease-related and other disease-related gene transcripts in blood

INVENTOR(S): Liew, Choong-Chin

PATENT ASSIGNEE(S): Chondrogene Limited, Can

SOURCE: U.S. Pat. Appl. Publ., 1/55 pp., Cont.-in-part of U.S.

Ser. No. 802,875. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE	
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PRIORITY APPLN. INFO.:			US 1999-115125P	P	19990106	<
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			US 2002-268730	A2	20021009	
			US 2003-601518	A2	20030620	
			US 2004-802875	A2	20040312	
			US 2004-812764	Α	20040330	

AB The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring diseases using gene-specific and/or tissue-specific primers. Affymetrix Human Genome U133 and ChondroChip microarrays were

used to detect differentially expressed gene transcripts in hypertension, obesity, allergy, systemic steroids, coronary artery disease, diabetes type 2, hyperlipidemia, lung disease, bladder cancer, rheumatoid arthritis, osteoarthritis, liver cancer, schizophrenia, Chagas disease, asthma, and manic depression syndrome. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen.

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L36 ANSWER 9 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN
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ACCESSION NUMBER:

2005:60754 HCAPLUS

DOCUMENT NUMBER:

Correction of: 2004:1036571 142:233342

Correction of: 142:16836

TITLE:

Sequences of human schizophrenia related genes and use

for diagnosis, prognosis and therapy

INVENTOR(S):

Liew, Choong-Chin

PATENT ASSIGNEE(S):

Chondrogene Limited, Can.

SOURCE:

U.S. Pat. Appl. Publ., 15 pp., Cont.-in-part of U.S.

Ser. No. 802,875.

CODEN: USXXCO

DOCUMENT TYPE:

Patent English

LANGUAGE: Eng FAMILY ACC. NUM. COUNT: 46

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
			- <i>f</i>		
US 2004241727	A1	20041202	u/s 2004-812731		20040330 <
US 2004014059	A1	20040122	∜ S 2002-268730		20021009 <
US 2005191637	A1	20050901	US 2004-803737		20040318 <
US 2005196762	A1	20050908	/US 2004-803759		20040318 <
US 2005196763	A1	20050908	US 2004-803857		20040318 <
US 2005196764	A1	20050908	US 2004-803858		20040318 <
US 2004241727	A1	20041202	US 2004-812731		20040330 <
US 2004241727	A1	20041202/	US 2004-812731		20040330 <
US 2004265869	A1	20041230	US 2004-812716		20040330 <
PRIORITY APPLN. INFO.:			US 1999-115125P	P	19990106 <
		/	US 2000-477148	В1	20000104
		/	US 2002-268730	A2	20021009
		<i>f</i>	US 2003-601518	A2	20030620
		/	US 2004-802875	A2	20040312
		/	US 2004-812731	A	20040330

The present invention is directed to detection and measurement of gene transcripts and their equivalent nucleic acid products in blood. Specifically provided is anal. performed on a drop of blood for detecting, diagnosing and monitoring diseases using gene-specific and/or tissue-specific primers. The present invention also describes methods by which delineation of the sequence and/or quantitation of the expression levels of disease-specific genes allows for an immediate and accurate diagnostic/prognostic test for disease or to assess the effect of a particular treatment regimen. [This abstract record is one of 3 records for this document necessitated by the large number of index entries required to fully index the document and publication system constraints.].

L36 ANSWER 10 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003: 7,96242 HCAPLUS

DOCUMENT NUMBER: 139:302975

Searched by Mary Jane Ruhl Ext. 22524

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Regulated expression systems for identification,
TITLE:
                                   screening, and directed synthesis of stabilized
                                   bioactive peptides for therapeutic use
                                   Altman, Elliot
INVENTOR(S):
                                   The University of Georgia Research Foundation, Inc,
PATENT ASSIGNEE(S):
                                   USA
SOURCE:
                                   U.S. Pat. Appl. Publ., 68 pp., Cont.-in-part of U.S.
                                   Ser. No. 701,947.
                                   CODEN: USXXCO
DOCUMENT TYPE:
                                   Patent
                                   English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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                                                             US 2002-210023
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       WO 2000022112
                                    A1
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       US 6818611
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       CA 2493306
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       WO 2004011485
                                             20040205
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                                    A2
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       WO 2004011485
                                             20050414
                                    A3
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RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
       EP 1537137
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                                    A2
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                 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                  IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
PRIORITY APPLN. INFO.:
                                                             US 1998-104013P
                                                                                       P 19981013 <--
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                                                             WO 1999-US23731
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                                                             US 2000-701947
                                                                                         A2 20001205
                                                             EP 1999-951940
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                                                             US 2002-210023
                                                                                         A 20020731
                                                                                         W 20030730
                                                             WO 2003-US23875
      An intracellular selecti\phin system allows screening for peptide bioactivity
AB
       and stability. Randomized recombinant peptides are screened for
       bioactivity in a tightly regulated expression system, preferably derived
       from the wild-type lac operon. Bioactive peptides thus identified are inherently protease- and peptidase-resistant. Also provided are bioactive
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peptides stabilized by a stabilizing group at the N-terminus, the C-terminus, or both. The stabilizing group can be a small stable protein, such as the Rop protein, **glutathione** sulfotransferase, **thioredoxin**, maltose binding protein, or **glutathione** reductase, an α -helical moiety, or one or more proline residues. Construction and characterization of a highly regulatable expression vector, pLAC11, and its multipurpose derivs., pLAC22 and pLAC33, is described. An in vivo approach for generating novel bioactive peptides that inhibit the growth of E. coli is disclosed. Directed synthesis of stable synthetically engineered inhibitor peptides is described.

L36 ANSWER 11 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:373944 HCAPLUS

DOCUMENT NUMBER: 133:118600

TITLE: Liver immunity and glutathione

AUTHOR(S): Yamauchi, Akira; Tsuyuki, Shigeru; Inamoto, Takashi;

Yamaoka, Yoshio

CORPORATE SOURCE: Department of Gastroenterological Surgery, Graduate

School of Medicine, Kyoto University, Kyoto, 606-8507,

Japan

SOURCE: Antioxidants & Redox Signaling (1999), 1(2),

245-253

CODEN: ARSIF2; ISSN: 1523-0864

PUBLISHER: Mary Ann Liebert

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review with 49 refs. Redox processes have been implicated in AB various biol. processes, including signal transduction, gene expression, and cell proliferation, and several mols. have been identified as redox regulators in cell activation. Glutathione is the oldest and most investigated mol. among them. Although details of the mechanisms by which glutathione regulates various aspects of cell biol. remains to be characterized, the relationship between immunodeficiency and cellular glutathione status is well established. Redox dysregulation contributes to the pathogenesis of acquired immunodeficiency syndrome (AIDS). Human immunodeficiency virus (HIV)-infected patients and simian immunodeficiency virus (SIV)-infected rhesus macaques have, on the average, significantly decreased plasma cysteine and intracellular glutathione levels. Liver contains abundant levels of reducing factors. However, glutathione levels in serum and peripheral blood mononuclear cells of cirrhosis patients are lower compared to values detected in healthy

individuals. In the present article, the significance of glutathione in regulating the functions of lymphocytes, especially those of liver-associated lymphocytes, has been described. A novel strategy for immune therapy of liver neoplasms with the use of redox

-modulating agents has been proposed.

REFERENCE COUNT: 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 12 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:312371 HCAPLUS

DOCUMENT NUMBER: 133:100789

TITLE: Modulation of DNA repair and glutathione

levels in human keratinocytes by micromolar arsenite

AUTHOR(S): Snow, Elizabeth T.; Hu, Yu; Yan, Chong Chao;

Chouchane, Salem

CORPORATE SOURCE: Nelson Institute of Environmental Medicine, New York

University School of Medicine, Tuxedo, NY, 10987, USA

Arsenic Exposure and Health Effects, Proceedings of SOURCE:

the International Conference on Arsenic Exposure and Health Effects, 3rd, San Diego, July 12-15, 1998 (

Chappell, Willard R.; Abernathy, Charles O.; Calderon,

1999), Meeting Date 1998, 243-251. Editor(s):

Rebecca L. Elsevier Science Ltd.: Oxford, UK.

CODEN: 68YOAM

DOCUMENT TYPE: Conference LANGUAGE: English

Arsenic (As) is a human carcinogen, but not a mutagen, although it inhibits DNA repair and is a comutagen. Human AG06 keratinocytes treated with micromolar arsenic exhibit dose and time-dependent loss of DNA ligase function. However, purified human DNA ligase I, ligase III, and other repair enzymes such as DNA polymerase β , are not inhibited by less than millimolar arsenite, As(III), the most toxic form of As found in the environment. DNA ligase activity in exts. from untreated keratinocytes is also insensitive to less than millimolar As. Pyruvate dehydrogenase, on the other hand, is inhibited by micromolar As and probably dets. As-induced cytotoxicity. Simultaneous treatment of AG06 cells with an alkylating agent, 1-methyl-3-nitro-1-nitrosoguanidine (MNNG), plus As produces a synergistic increase in viability (dye uptake) at low doses and a synergistic increase in toxicity at high doses. Micromolar As also modulates cellular redox levels and induces a variety of cellular stress response genes. Keratinocytes treated with As exhibit both a time- and dose-dependent increase in cellular GSH levels and alterations in the relative activity of several GSH-dependent enzymes. These As-induced changes in cellular redox capacity and DNA repair activity are not directly related to toxicity. Maximal induction of GSH and DNA repair occurs after treatment with sub-toxic concns. of As. At submicromolar concns., arsenic also induces hyperproliferation of keratinocytes, both in vivo and in vitro. These results suggest that As modulates DNA repair and redox levels primarily through post-translational or transcriptional mechanisms.

REFERENCE COUNT: THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS 4.3 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 13 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:141601 HCAPLUS

Correction of: 1996:263007

DOCUMENT NUMBER: 132:150093

Correction of: 124:314095

TITLE: Reduction-oxidation (redox) state regulation

of extracellular matrix metalloproteinases and tissue

inhibitors in cardiac normal and transformed

fibroblast cells

AUTHOR (S): Tyagi, Suresh C.; Kumar, G. Suresh; Broders, Susan CORPORATE SOURCE: Dalton Cardiovascular Research Center, University

Missouri-Columbia, Columbia, MO, 65212, USA

SOURCE: Journal of Cellular Biochemistry (1996),

61(1), 139-151

CODEN: JCEBD5; ISSN: 0730-2312

PUBLISHER: Wiley-Liss DOCUMENT TYPE: Journal LANGUAGE: English

Latent matrix metalloproteinases (MMPs) in normal myocardium are activated in end-stage heart failure. In vitro oxidized glutathione

(GSSG) activates myocardial MMPs which contains a cysteine residue. In vivo GSSG induce the collagen lysis and cardiac dilatation. To assess

whether thiol and non-thiol reducing agents have

direct effect on the interstitial human heart fibroblast (HHF) proliferation and MMP expression, HHF and polyoma virus transformed fibroblast cells were cultured with or without the thiol-containing reduced (GSH) or oxidized (GSSG) glutathiones , pyrrolidine dithiocarbamate (PDTC) and N-acetylcysteilne (NAC), and non-thiol ascorbic acid. After 100 μq/mL (.apprx.0.3 mM) GSH or PDTC treatment the proliferative (synthetic) phenotype of transformed fibroblast cells was changed to quiescent (contractile) phenotype. Also, after GSH, PDTC, and ascorbic acid treatment the medium was then analyzed for MMP activity by zymog. The results indicate reduction in MMP expression in transformed fibroblast cells after GSH and PDTC treatments and no effect after ascorbic acid treatment. Based on reverse zymog., we observed the level of tissue inhibitor of metalloproteinase (TIMP) at a decreased level in transformed cells. The effect of the reducing agent at the gene transcription was measured by estimating mRNA (Northern blot anal.) of MMP and of TIMP in the cells that were cultured in medium in the presence and absence of GSH. These results indicate that GSH induces MMP-2 and MMP-1 expression in normal HHF and that GSH reduces MMP-2 and MMP-1 in transformed fibroblast cells. After the treatment, the TIMP-2 level was repressed in normal HHF and TIMP-2 level increased in transformed fibroblast cells. These events are dependent on the nuclear transcription factor activity on the collagenase promoter in normal HHF cells. On the other hand, in polyoma transform fibroblast cells these events are not dependent on this collagenase promoter. These results suggest that oxidative environment induces normal HHF cell proliferation, and the reducing agent decreases normal HHF cell proliferation by inducing MMP and repressing TIMP gene transcription. In transformed cells reducing agents inhibit MMP expression and increase TIMP levels, which suggests a role of antioxidants in preventing tumorigenesis.

L36 ANSWER 14 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:98240 HCAPLUS

DOCUMENT NUMBER: 132:146655

TITLE: Inhibitors of redox signaling for

restoration of apoptosis and inhibition of abnormal

cell proliferation

INVENTOR(S): Kirkpatrick, D. Lynn; Powis, Garth

PATENT ASSIGNEE(S): Can.

SOURCE: PCT Int. Appl., 55 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		DATE				
WO 2000 WO 2000				A2 A3		20000210 20000615			WO 1	 999-1	US17	-	19990802 <					
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	TM,		TT,	UA,		PL, US,												
RW:	ES,	FI,	FR,	GB,	GR,	SD, IE, ML,	IT,	LU,	MC,	NL,	PT,							

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    US 6372772
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                        A1
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                                         US 1997-54566P
PRIORITY APPLN. INFO.:
                                                            A 19980731 <--
                                         US 1998-127219
                                          WO 1999-US17496
                                                             W 19990802
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AB The present invention is directed to a composition or formulation which inhibits or interferes with cellular redox function, and method of using same to restore normal cellular function. More specifically, the composition of the present invention interferes with or inhibits abnormal cellular proliferation and or restores cellular apoptosis.

L36 ANSWER 15 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:778665 HCAPLUS

DOCUMENT NUMBER: 132:147086

TITLE: Increased glutathione synthesis associated

with platelet-derived growth factor stimulation of

NIH3T3 fibroblasts

AUTHOR(S): Iantomasi, T.; Favilli, F.; Degl'Innocenti, D.;

Vincenzini, M. T.

CORPORATE SOURCE: viale Morgagni 50, Department of Biochemical Sciences,

University of Florence, Florence, 50134, Italy

SOURCE: Biochimica et Biophysica Acta, Molecular Cell Research

(1999), 1452(3), 303-312

CODEN: BBAMCO; ISSN: 0167-4889

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

ΔR Previous data show a relation between GSH content and proliferation of normal and tumor cells. We recently demonstrated a specific involvement of GSH in the autophosphorylation activity of the platelet-derived growth factor (PDGF) receptor in NIH3T3 fibroblasts. In this study we demonstrate that the stimulation by PDGF of serum-starved NIH3T3 cells increases cellular GSH content, while no change in oxidized GSH content was measured. Expts. performed with actinomycin, cycloheximide and buthionine sulfoximide, a specific inhibitor of the rate-limiting enzyme of the de novo synthesis of GSH γ glutamylcysteine synthetase $(\gamma\text{-GCS})$, confirm PDGF induction of GSH synthesis. These results provide the first demonstration that PDGF mediated transduction signals seem strictly related to mechanisms involved in the increase of γ -GCS activity associated with increased γ -GCS heavy subunit mRNA levels. In fact, serum and epidermal growth factor (EGF) stimulation of quiescent NIH3T3 and NIH3T3, which overexpress EGF receptor, does not affect GSH content or its synthesis. These data may be related to a possible GSH role in the redox regulation of cell proliferation mediated by PDGF.

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 16 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:582008 HCAPLUS

DOCUMENT NUMBER: 131:333310

TITLE: Effect of Thioacetamide on the Hepatic Expression of

 γ -Glutamylcysteine Synthetase Subunits in the

Rat

AUTHOR(S): Lu, Shelly C.; Huang, Zong-Zhi; Yang, Heping;

Tsukamoto, Hidekazu

CORPORATE SOURCE: Division of Gastroenterology and Liver Diseases, USC

Liver Disease Research Center, USC School of Medicine,

Los Angeles, CA, 90033, USA

SOURCE: Toxicology and Applied Pharmacology (1999),

159(3), 161-168

CODEN: TXAPA9; ISSN: 0041-008X

PUBLISHER: Academic Press

DOCUMENT TYPE: Journal LANGUAGE: English

AB Glutathione (GSH) is the main non-protein thiol

important in antioxidant defense and maintenance of the intracellular redox state. A major determinant of the rate of GSH synthesis is the activity of the rate-limiting enzyme, γ-glutamylcysteine synthetase (GCS). A heavy (HS) and light subunit (LS) make up GCS; oxidative stress regulates both transcriptionally. cis-Acting elements important for the oxidative stress-induced transcriptional up-regulation of both subunits are antioxidant response element (ARE) and activator protein-1 (AP-1) site. The nuclear factor-κB (NF-κB) binding site may also regulate the heavy subunit. Increased GSH and γ-glutamyltranspeptidase are often observed in preneoplastic hepatocyte nodules and may be important in hepatocarcinogenesis. The current work examined the effect of a commonly used hepatocarcinogen, thioacetamide (TAA), on the expression of GCS subunits. After 3 wk of TAA treatment, liver GSH level remained unchanged despite significant oxidative stress as measured by the thiobarbituric acid reactive substance assay. The mRNA levels of GCS-HS and GCS-LS increased six- and fourfold, resp., and the protein level of GCS-HS and GCS activity all increased. Electrophorectic mobility shift assay showed binding to ARE, AP-1, and NF-kB probes all increased. These results suggest TAA treatment increased hepatic GCS subunit expression and GCS activity by inducing oxidative stress and increasing the binding to redox-sensitive cis-acting elements important for transcriptional up-regulation of GCS. This is the first in vivo study that examined the effect of a hepatocarcinogen on GCS expression. (c) 1999 Academic Press.

REFERENCE COUNT:

54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 17 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:569467 HCAPLUS

DOCUMENT NUMBER: 131:320870

TITLE: Redox modulation of cell surface protein

thiols in U937 lymphoma cells: the role of γ -glutamyl transpeptidase-dependent H2O2

production and S-thiolation

AUTHOR(S): Dominici, S.; Valentini, M.; Maellaro, E.; Del Bello,

B.; Paolicchi, A.; Lorenzini, E.; Tongiani, R.;

Comporti, M.; Pompella, A.

CORPORATE SOURCE: Institute of General Pathology, University of Siena,

Siena, Italy

SOURCE: Free Radical Biology & Medicine (1999),

27(5/6), 623-635

CODEN: FRBMEH; ISSN: 0891-5849

PUBLISHER: Elsevier Science Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

AB The expression of gamma-glutamyl transpeptidase (GGT), a plasma membrane ectoenzyme involved in the metabolism of extracellular reduced

glutathione (GSH), is a marker of neoplastic progression in several exptl. models, and occurs in a number of human malignant neoplasms and their metastases. Because it favors the supply of

precursors for the synthesis of GSH, GGT expression has been interpreted as a member in cellular antioxidant defense systems. However, thiol metabolites generated at the cell surface during GGT activity can induce prooxidant reactions, leading to production of free radical oxidant species. The present study was designed to characterize the prooxidant reactions occurring during GGT ectoactivity, and their possible effects on the thiol redox status of proteins of the cell surface. Results indicate that: (i) in U937 cells, expressing significant amts. of membrane-bound GGT, GGT-mediated metabolism of GSH is coupled with the extracellular production of hydrogen peroxide; (ii) GGT activity also results in decreased levels of protein thiols at the cell surface; (iii) GGT-dependent decrease in protein thiols is due to sulfhydryl oxidation and protein S-thiolation reactions; and (iv) GGT irreversible inhibition by acivicin is sufficient to produce an increase of protein thiols at the cell surface. Membrane receptors and transcription factors have been shown to possess critical thiols involved in the transduction of proliferative signals. Furthermore, it was suggested that Sthiolation of cellular proteins may represent a mechanism for protection of vulnerable thiols against irreversible damage by prooxidant agents. Thus, the findings reported here provide addnl. explanations for the envisaged role played by membrane-bound GGT activity in the proliferative attitude of malignant cells and their resistance to prooxidant drugs and radiation therapy.

REFERENCE COUNT:

60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 18 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1999:467538 HCAPLUS

DOCUMENT NUMBER:

131:241212

TITLE:

Differential reconstitution of mitochondrial respiratory chain activity and plasma **redox** state by cysteine and ornithine in a model of

cancer cachexia

AUTHOR (S):

CORPORATE SOURCE:

Ushmorov, Alexej; Hack, Volker; Droge, Wulf Deutsches Krebsforschungszentrum, Division of Immunochemistry, Heidelberg, D-69120, Germany

SOURCE: Cancer Research (1999), 59(14), 3527-3534

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER:

AACR Subscription Office

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The mechanism of wasting, as it occurs in malignant diseases and various AB etiol. unrelated conditions, is still poorly understood. The authors have, therefore, studied putative cause/effect relationships in a murine model of cancer cachexia, C57BL/6 mice bearing the fibrosarcoma MCA-105. The plasma of these mice showed decreased albumin and increased glutamate levels, which are typically found in practically all catabolic conditions. Skeletal muscles from tumor-bearing mice were found to have an abnormally low mitochondrial respiratory chain activity (mito.RCA) and significantly decreased glutathione (GSH) levels. The decrease in mito.RCA was correlated with an increase in the i.m. GSH disulfide/GSH ratio, the plasma cystine/thiol ratio, and the GSH disulfide/GSH ratio in the bile. This is indicative of a generalized shift in the redox state extending through different body fluids. Treatment of tumor-bearing mice with ornithine, a precursor of the radical scavenger spermine, reversed both the decrease in mito.RCA and the change in the redox state, whereas treatment with cysteine, a GSH precursor, normalized only the redox state.

Treatment of normal mice with difluoromethyl-ornithine, a specific inhibitor of ornithine decarboxylase and spermine biosynthesis, inhibited the mito.RCA in the skeletal muscle tissue, thus illustrating the importance of the putrescine/spermine pathway in the maintenance of mito.RCA. Ornithine, cysteine, and N-acetyl-cysteine (NAC) also reconstituted the abnormally low concns. of the GSH precursor glutamate in the skeletal muscle tissue of tumor-bearing mice. Higher doses, however, enhanced tumor growth and increased the plasma glucose level in normal mice. In the latter, cysteine and NAC also decreased i.m. catalase and GSH peroxidase activities. Taken together, the studies on the effects of ornithine, cysteine, and NAC illuminate some of the mechanistic pathways involved in cachexia and suggest targets for therapeutic intervention.

THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 51 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 19 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:25288 HCAPLUS

DOCUMENT NUMBER: 130:195112

TITLE: Changes in glutathione status and the

antioxidant system in blood and in cancer cells associate with tumor growth in vivo

AUTHOR(S): Navarro, Jose; Obrador, Elena; Carretero, Julian;

Petschen, Ignacio; Avino, Jose; Perez, Pilar; Estrela,

Jose M.

CORPORATE SOURCE: Departamento de Fisiologia, Universidad de Valencia,

Facultad de Medicina, Valencia, 46010, Spain

SOURCE: Free Radical Biology & Medicine (1998),

Volume Date 1999, 26(3/4), 410-418 CODEN: FRBMEH; ISSN: 0891-5849

PUBLISHER: Elsevier Science Inc.

Journal DOCUMENT TYPE: LANGUAGE:

humans.

English The relationship among cancer growth, the glutathione

redox cycle and the antioxidant system was studied in blood and in

tumor cells. During cancer growth, the glutathione redox status (GSH/GSSG) decreases in blood of Ehrlich ascites tumor-bearing mice. This effect is mainly due to an increase in GSSG levels. Two reasons may explain the increase in blood GSSG: (a) the increase in peroxide production by the tumor that, in addition to changes affecting the glutathione-related and the antioxidant enzyme activities, can lead to GSH oxidation within the red blood cells; and (b) an increase of GSSG release from different tissues into the blood. GSH and peroxide levels are higher in the tumor cells when they proliferate actively, however GSSG levels remain constant during tumor growth in mice. These changes associate with low levels of lipid peroxidn. in plasma, blood and the tumor cells. The GSH/GSSG ratio in blood also decreases in patients bearing breast or colon cancers and, as it occurs in tumor -bearing mice, this change assocs. with higher GSSG levels, especially in advanced stages of cancer progression. Our results indicate that determination of glutathione status and oxidative stress-related parameters in blood may help to orientate cancer therapy in

REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 20 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN ACCESSION NUMBER: 1998:400616 HCAPLUS

DOCUMENT NUMBER: 129:134396

TITLE: Disruption of redox homeostasis in the

transforming growth factor- α/c -myc transgenic

mouse model of accelerated

hepatocarcinogenesis

AUTHOR(S): Factor, Valentina M.; Kiss, Andras; Woitach, Joseph

T.; Wirth, Peter J.; Thorgeirsson, Snorri S.

CORPORATE SOURCE: Laboratory of Experimental Carcinogenesis, NCI,

National Institutes of Health, Bethesda, MD, 20892,

USA

SOURCE: Journal of Biological Chemistry (1998),

273 (25), 15846-15853

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER: American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE: Journal LANGUAGE: English

AB In previous studies the authors have demonstrated that transforming growth

factor (TGF)- α/c -myc double transgenic mice exhibit an enhanced rate

of cell **proliferation**, accumulate extensive DNA damage, and develop multiple liver **tumors** between 4 and 8 mo of age. To

clarify the biochem. events that may be responsible for the genotoxic and

carcinogenic effects observed in this transgenic model, several parameters of redox homeostasis in the liver were examined prior to development of hepatic tumors. By 2 mo of age, production of

reactive oxygen species, determined by the peroxidn.-sensitive fluorescent dye,

2',7'-dichlorofluorescein diacetate, was significantly elevated in TGF- α/c -myc transgenic hepatocytes vs. either wild type or c-myc single transgenic cells, and occurred in parallel with an increase in lipid peroxidn. Concomitantly with a rise in oxidant levels, antioxidant

defenses were decreased, including total glutathione content and

the activity of glutathione peroxidase, whereas

thioredoxin reductase activity was not changed. However, hepatic

tumors which developed in $TGF-\alpha/c$ -myc mice exhibited an increase in thioredoxin reductase activity and a very low activity of glutathione peroxidase. Furthermore, specific

deletions were detected in mtDNA as early as 5 wk of age in the transgenic

mice. These data provide exptl. evidence that co-expression of TGF- α and c-myc transgenes in mouse liver promotes overprodn. of

reactive oxygen species and thus creates an oxidative stress environment. This phenomenon may account for the massive DNA damage and acceleration of

hepatocarcinogenesis observed in the TGF- α /c-myc mouse model.

REFERENCE COUNT: 81 THERE ARE 81 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 21 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:369893 HCAPLUS

DOCUMENT NUMBER: 129:89983

TITLE: Antioxidants reduce cyclooxygenase-2 expression,

prostaglandin production, and proliferation

in colorectal cancer cells

AUTHOR(S): Chinery, Rebecca; Beauchamp, R. Daniel; Shyr, Yu;

Kirkland, Susan C.; Coffey, Robert J.; Morrow, Jason

D.

CORPORATE SOURCE: Department of Medicine, The Vanderbilt Cancer Center,

Vanderbilt University Medical Center, Nashville, TN,

37232, USA

SOURCE: Cancer Research (1998), 58(11), 2323-2327

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

AB Increased expression of cyclooxygenase (COX) and overprodn. of prostaglandins (PGs) have been implicated in the development and progression of colorectal cancer (CRC). Recent observations suggest that reactive oxygen intermediates play a role in tumor cell growth regulation and expression of the inducible COX, COX-2. Therefore, the effects of various antioxidants on COX expression and cellular growth were evaluated in the human CRC cell line HCA-7. The antioxidants pyrrolidinedithiocarbamate (PDTC), N-acetylcysteine, 6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid (Trolox), and U74006 decreased PG production, intracellular redox status, and cellular growth in a concentration-dependent manner. The decrease

in

cellular growth was associated with the induction of apoptosis. Unlike the selective COX inhibitors 1-[(4-methylsulfonyl)phenyl]-3-trifluoromethyl-5-[(4-fluoro)phenyl]pyrazole (SC 58125) and (2-cyclohexyloxy-4-nitrophenyl)methanesulfonamide (NS 398) that inhibit COX-2 catalytic activity, these antioxidants decreased COX-2 expression at the transcriptional level. Combined treatment of HCA-7 cells with PDTC and SC 58125 resulted in an additive decrease in PG levels and anchorage-dependent and -independent growth. Furthermore, whereas antioxidants or SC 58125 reduced tumor growth in vivo, coadministration of PDTC and SC 58125 resulted in actual tumor regression. These results suggest that combined therapy with NSAIDs and antioxidants might be useful in the prevention and/or treatment of CRC.

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 22 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:229894 HCAPLUS

DOCUMENT NUMBER: 129:234

TITLE: Mechanisms of inhibition of the thioredoxin

growth factor system by antitumor

2-imidazolyl disulfides

AUTHOR(S): Kirkpatrick, D. Lynn; Kuperus, Miles; Dowdeswell,

Marla; Potier, Noelle; Donald, Lynda J.; Kunkel, Mark;

Berggren, Margareta; Angulo, Miguel; Powis, Garth

CORPORATE SOURCE: Department of Chemistry, University of Regina, Regina,

SK, S4S 0A2, Can.

SOURCE: Biochemical Pharmacology (1998), 55(7),

987-994

CODEN: BCPCA6; ISSN: 0006-2952

PUBLISHER: Elsevier Science Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

The interactions of a series of 2-imidazolyl disulfide antitumor compds. with the thioredoxin reductase (TR)-thioredoxin (hTrx) redox system have been studied. Bu 2-imidazolyl disulfide (I) and Et 2-imidazolyl disulfide (II) were substrates for reduction by TR with Km values of 43 and 48 µM. 1-Methylpropyl 2-imidazolyl disulfide (III) and benzyl 2-imidazolyl disulfide (IV) were competitive inhibitors of the reduction of hTrx by TR with Ki values of 31 µM. None of the disulfides were substrates for reduction by human glutathione reductase. The disulfides caused reversible thioalkylation of hTrx at the redox catalytic site as shown by the fact that there was no thioalkylation of a mutant hTrx where both the catalytic site Cys32 and Cys35 residues were replaced by Ser. In addition, the disulfides caused a

slower irreversible inactivation of hTrx as a substrate for reduction by TR, with half-lives for I of 30 min, for III of 4 h, and for tert-Bu 2-imidazolyl disulfide of 24 h. This irreversible inactivation of hTrx occurred at concns. of the disulfides an order of magnitude below those that inhibited TR, and involved the Cys73 of hTrx, which is outside the conserved redox catalytic site, as shown by the resistance to inactivation of a mutant hTrx where Cys73 was replaced by Ser. Electrophoretic and mass spectral analyses of the products of the reaction between the disulfides and hTrx show that modification of 1-3 Cys residues of the protein occurred in a concentration-dependent fashion. The disulfides inhibited the hTrx-dependent proliferation of MCF-7 breast cancer cells with IC50 values of I and III of 0.2 and 1.2 μM, resp. The results show that although the catalytic sites of TR and hTrx are reversibly inhibited by the 2-imidazolyl disulfides, it is the irreversible thioalkylation of Cys73 of hTrx by the disulfides that most probably accounts for the inhibition of thioredoxin-dependent cell grown by the disulfides.

REFERENCE COUNT:

26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 23 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1998:163492 HCAPLUS

DOCUMENT NUMBER:

128:213410

TITLE:

Modulators of nitrosative and oxidative stress for the

treatment of disease

INVENTOR(S):

Stamler, Jonathan S.; Griffith, Owen W.

PATENT ASSIGNEE(S):

Duke University, USA; Medical College of Wisconsin

Research Foundation, Inc.

SOURCE:

PCT Int. Appl., 159 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9808566 W: AU, CA, JP	A1 19980305	WO 1997-US13876	19970813 <
RW: AT, BE, CH,	DE, DK, ES, FI,	FR, GB, GR, IE, IT, I	LU, MC, NL, PT, SE
US 6057367	A 20000502	US 1997-852490	19970507 <
CA 2262708	AA 19980305	CA 1997-2262708	19970813 <
AU 9740542		AU 1997-40542	19970813 <
EP 963219	A1 19991215	EP 1997-938149	19970813 <
R: CH, DE, ES,	FR, GB, IT, LI,	NL, SE	
US 6180824	B1 20010130	US 1999-361167	19990727 <
US 6359004	B1 20020319	US 2000-690989	20001018 <
US 2003096870	A1 20030522	US 2001-13455	20011213 <
US 6608110	B2 20030819		
US 2003207815	A1 20031106	US 2003-417238	20030417 <
PRIORITY APPLN. INFO.:		US 1996-25819P	P 19960830 <
		US 1997-852490	A 19970507 <
		WO 1997-US13876	W 19970813 <
		US 1999-361167	A1 19990727
		US 2000-690989	A1 20001018
		US 2001-13455	A3 20011213

AB Mammals are treated for infections or for conditions associated with pathol. proliferating mammalian cell growth (for example, certain cancers, restenosis, benign prostatic hypertrophy) by

administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathol. proliferating mammalian cells. Novel agents include α -alkyl-homocysteine sulfoximines wherein the α -alkyl contains 2-8 carbon atoms, and the S-alkyl contains 1-10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g. humans at risk for a stroke because of having had a transient ischemic attack are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 24 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:84114 HCAPLUS

DOCUMENT NUMBER: 128:179261

TITLE: Thiol redox modulation of

tumor necrosis factor- α responsiveness

in cultured AIDS-related Kaposi's sarcoma cells

AUTHOR(S): Mallery, S. R.; Landwehr, D. J.; Ness, G. M.; Clark,

Y. M.; Hohl, C. M.

CORPORATE SOURCE: Departments of Oral Surgery and Pathology, Colleges of

Dentistry and Medicine, Ohio State University,

Columbus, OH, 43210, USA

SOURCE: Journal of Cellular Biochemistry (1998),

68(3), 339-354

CODEN: JCEBD5; ISSN: 0730-2312

PUBLISHER: Wiley-Liss, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

Both clin. and exptl. evidence indicates that AIDS-related Kaposi's sarcoma (AIDS-KS) has a multifactorial pathogenesis with factors such as HIV viral load, latent virus induction, and opportunistic infections contributing to disease progression. However, a consistent feature that unites these apparently diverse putative etiol. agents is sustained serum elevations of pro-inflammatory cytokines such as tumor necrosis factor- α (TNF- α). While virtually every cell responds to $TNF-\alpha$ with gene activation, the extent of $TNF-\alpha$ -mediated cellular signaling is regulated by a delicate balance between signal activation and signal arresting events. Reactive oxygen intermediates (ROI), which are generated as a consequence of TNF- α membrane interaction, are part of this $TNF-\alpha$ -initiated cellular activation cascade. Previous studies in the authors' laboratory have shown that AIDS-KS cells possess impaired oxygen intermediate scavenging capacities, thereby establishing conditions permissive for the intracellular retention of ROI. Here, the authors used cellular capacity to upregulate the cytoprotective enzyme superoxide dismutase (SOD) to address the extent of cellular response to $TNF-\alpha$. Concurrent with the SOD analyses, nucleotide profiles were obtained to assess cellular bioenergetic responses during $TNF-\alpha$ challenge. **Proliferative** growth levels of mitochondrial (Mn)SOD activities showed an activity spectrum ranging from lowest activity in AIDS-KS cells, to intermediate levels in matched, nonlesional cells from the AIDS-KS donors, to highest activities in HIV normal fibroblasts. In contrast, following TNF- α challenge, the

AIDS-KS and KS donor nonlesional cells showed a 11.89- and 5.86-fold resp. increase in MnSOD activity, while the normal fibroblasts demonstrated a 1.35-fold decrease. Subsequent thiol redox modulation studies showed that only the normal fibroblast cultures showed a potentiation of $TNF-\alpha$ -mediated MnSOD upregulation following GSH depletion. In addition, provision of the GSH precursor, Nacetylcysteine during TNF- α challenge only diminished MnSOD activity and mitochondrial compartmentalization in the AIDS-KS cells, a finding that likely reflects the lower levels of reduced thiols in this cellular population. The authors' data, which show that a perturbation in their cellular thiol redox status accentuates AIDS-KS cellular responsiveness to TNF-α, suggest a biochem. rationale for the recognized TNF- α AIDS-KS clin. correlation.

REFERENCE COUNT:

51 THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 25 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1998:49693 HCAPLUS

DOCUMENT NUMBER:

128:149308

TITLE:

Cellular thioredoxin reductase activity is

regulated by selenium

AUTHOR (S):

Berggren, Margareta; Gallegos, Alfred; Gasdaska, John;

Powis, Garth

CORPORATE SOURCE:

Arizona Cancer Center, University of Arizona Health

Sciences Center, Tucson, AZ, 85724-5024, USA

SOURCE:

Anticancer Research (1997), 17(5A),

3377-3380

CODEN: ANTRD4; ISSN: 0250-7005

PUBLISHER:

Anticancer Research

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Selenium (Se) is an essential trace element and has been reported to AB decrease the incidence of some human cancers. The authors have investigated the effects of Se on thioredoxin reductase, a selenocysteine containing flavoenzyme, in HT-29 human colon cancer cells grown in serum-free medium. Sodium selenite and other Se containing compds. produced a time and concentration dependent increase in intracellular thioredoxin reductase activity and protein levels. Selenite was the most active of the Se compds. examined: 1 μM selenite produced a 28-fold increase in thioredoxin reductase activity by 1 day and 10 µM selenite over a 60-fold increase by 5 days. The activity of a related non-selenocysteine containing flavoenzyme glutathione reductase was not increased by selenite. Selenite, but not the other Se containing compds. inhibited cell growth at concns. above 2 µM. results show that Se can produce large increases in cell thioredoxin reductase activity.

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS 30 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 26 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1997:729411 HCAPLUS

DOCUMENT NUMBER:

128:21764

TITLE:

Role of intracellular redox status in

apoptosis induction of human T-cell leukemia virus type I-infected lymphocytes by 13-cis-retinoic acid

AUTHOR (S):

Furuke, Keizo; Sasada, Tetsuro; Ueda-Taniguchi, Yasuyo; Yamauchi, Akira; Inamoto, Takashi; Yamaoka,

Yoshio; Masutani, Hiroshi; Yodoi, Junji

CORPORATE SOURCE:

Department of Biological Responses, Institute for

Virus Research, Kyoto University, Kyoto, 606-01, Japan

SOURCE:

Cancer Research (1997), 57(21), 4916-4923

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER:

American Association for Cancer Research

DOCUMENT TYPE:

Journal

LANGUAGE: English

We have shown that cell cycle progression of human T-cell leukemia virus type I (HTLV-I)-transformed T-cell lines was inhibited by 13-cis-retinoic acid (13cRA). In the present study, we report that 13cRA inhibited proliferation and induced cell death of peripheral blood mononuclear cells obtained from four patients with acute adult T-cell leukemia but not of mitogen- or interleukin 2-activated peripheral blood mononuclear cells from HTLV-I-neg. healthy donors. Because HTLV-I-infected lymphocytes are susceptible to oxidative stress, we examined the role of the intracellular redox state in 13cRA-induced cell death using a HTLV-I-pos. T-cell line, ATL2, as a model. The 13cRA induced apoptosis in ATL2 cells within 48 h in a dose-dependent manner. The ability of 13cRA to induce apoptosis was more potent than that of all-trans-retinoic acid. Apoptosis induction by 13cRA was significantly enhanced by buthionine sulfoximine (BSO), which decreased the levels of intracellular reduced glutathione, although 13cRA by itself did not alter them, suggesting that intracellular reduced glutathione may modulate 13cRA-induced apoptosis. In addition, flow cytometric anal. revealed that 13cRA increased intracellular peroxides in 24 h and that the addition of BSO further enhanced them. Although N-acetylcysteine had only a marginal effect, pretreatment with catalase markedly inhibited 13cRA-induced apoptosis. These results suggest that peroxide generation, i.e., oxidative stress, may play a crucial role in the induction of apoptosis by 13cRA and further demonstrate that combined treatment with 13cRA and BSO induces apoptosis of HTLV-I-pos. lymphocytes even more potently.

REFERENCE COUNT:

THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS 51 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 27 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1997:561551 HCAPLUS

DOCUMENT NUMBER:

127:218945

TITLE:

Nitric oxide and superoxide induced p53 and Bax

accumulation during mesangial cell apoptosis

AUTHOR(S):

Sandau, Katrin; Pfeilschifter, Josef; Brune, Bernhard

CORPORATE SOURCE: Faculty of Medicine, Department of Medicine IV,

Experimental Division, University of

Erlangen-Nurnberg, Erlangen, Germany

Kidney International (1997), 52(2), 378-386 CODEN: KDYIA5; ISSN: 0085-2538

PUBLISHER:

SOURCE:

Blackwell

DOCUMENT TYPE:

Journal

LANGUAGE:

English

During proliferative glomerulonephritis, the early phase of mesangiolysis is linked to increased nitric oxide (NO) production NO as well as superoxide (02-) are inflammatory mediators that are generated by mesangial cells (MC) after cytokine stimulation. Added individually, both radicals induce MC apoptosis. However, the coexistence of a defined NO/O2- ratio is cross-protective. Apoptosis is characterized by specific features such as chromatin condensation, DNA strand breaks, and the occurrence of apoptotic regulating proteins. The tumor suppressor p53 and Bax (Bcl-2 associated protein x) are considered to be classical death promoters, which accumulate after toxic insults. To study

p53 and Bax protein accumulation in NO and/or O2--induced apoptosis, the authors used the NO-donor S-nitrosoglutathione (GSNO) and the redox cycler 2,3-dimethoxy-1,4-naphthoquinone (DMNQ). Both agonists initiated DNA fragmentation in a concentration dependent manner associated

with transient p53 and Bax up-regulation. Co-generation of NO/O2resulted not only in reduced DNA fragmentation, but also in decreased Bax accumulation. Comparable to the NO/O2- co-generation, cytokines failed to induce apoptosis. In contrast, cytokines in combination with pyrrolidine dithiocarbamate, which blocks endogenous superoxide dismutase, allowed p53 and Bax accumulation as well as DNA fragmentation. results demonstrate p53 and Bax as early components in NO and O2--induced rat MC apoptosis and point to the NO/O2- interaction as a naturally occurring cell defense mechanism.

REFERENCE COUNT: 51 THERE ARE 51 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 28 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

1997:556476 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 127:275947

TITLE: Generation of angiostatin by reduction and proteolysis

of plasmin. Catalysis by a plasmin reductase secreted

by cultured cells

AUTHOR (S): Stathakis, Paul; Fitzgerald, Melinda; Matthias, Lisa

J.; Chesterman, Colin N.; Hogg, Philip J.

CORPORATE SOURCE: Centre for Thrombosis and Vascular Research, School of

Pathology and Department of Haematology, Prince of Wales Hospital, University of New South Wales, Sydney,

NSW 2052, Australia

Journal of Biological Chemistry (1997), SOURCE:

272(33), 20641-20645

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER: American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE: Journal LANGUAGE: English

Extracellular manipulation of protein disulfide bonds has been implied in diverse biol. processes, including penetration of viruses and endotoxin into cells and activation of certain cytokine receptors. We now demonstrate reduction of one or more disulfide bonds in the serine proteinase, plasmin, by a reductase secreted by Chinese hamster ovary or HT1080 cells. Reduction of plasmin disulfide bond(s) triggered proteolysis of the enzyme, generating fragments with the domain structure of the angiogenesis inhibitor, angiostatin. Two of the known reductases secreted by cultured cells are protein disulfide isomerase and thioredoxin, and incubation of plasmin with these purified reductases resulted in angiostatin fragments comparable with those generated from plasmin in cell culture. Thioredoxin-derived angiostatin inhibited proliferation of human dermal microvascular endothelial cells with half-maximal effect at approx. 0.2 $\mu g/mL$. Angiostatin made by cells and by purified reductases contained free sulfhydryl group(s), and S-carbamidomethylation of these thiol group(s) ablated biol. activity. Neither protein disulfide isomerase nor thioredoxin were the reductases used by cultured cells, because immunodepletion of conditioned medium of these proteins did not affect angiostatin generating activity. The plasmin reductase secreted by HT1080 cells required a small cofactor for activity, and physiol. relevant concns. of reduced glutathione fulfilled this role. These results have consequences for plasmin activity and angiogenesis, particularly in the context of

tumor growth and metastasis. Moreover, this is the first demonstration of extracellular reduction of a protein disulfide bond, which has general implications for cell biol.

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 29 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:511962 HCAPLUS

DOCUMENT NUMBER: 127:117382

TITLE: Oxidized glutathione, salts, and derivatives

as enhancers of endogenous production of cytokines and hemopoietic factors, and methods of therapeutic use

INVENTOR(S): Balazovsky, Mark Borisovich; Kozhemyakin, Leonid

Andreevich

Patent

PATENT ASSIGNEE(S): Balazovsky, Mark Borisovich, Russia; Kozhemyakin,

Leonid Andreevich

SOURCE: PCT Int. Appl., 125 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE: English FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PAT	CENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE		
WO	9721	444			A1		1997	0619		₩O 1	996-	R1134	Λ		1	9961	210	<i></i> -
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ΕP	8698	09			A1		1998	1014				9419						
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	R:	AT,	BE,	CH,	DE,	DK.	ES,	FR.	GB,	GR,	IT.	LI.	LU.	NL.	SE.	MC.	PT.	
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RU	2153	351			C2		2000	0727		RU 1	998-	1080	38		1	9961	210	<
JP	2000	5151	11		T2		2000	1114		JP 1	997-	5219	65		1	9961	210	<
ΑT	2149	36			E		2002	0415		AT 1	996-	9419	15		1	9961	210	<
US	6492	329			В1		2002	1210	1	US 2	000-	7027	01			0001		
RITY	APP	LN.	INFO	. :						RU 1	995-	1204	03		A 1	9951	214	<
									1	WO 1	996-1	RU22			A 1	9960	808	<
												7338				9961		
												RU34			A 1	9961	210	<
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US 1996-766557 A 19961211 <--

AB A method for stimulating endogenous production of cytokines and hemopoietic factors comprises topical or parenteral administration of an effective amount of oxidized glutathione, and/or a pharmaceutically acceptable salt and/or derivative thereof, for a period sufficient to stimulate the endogenous production to obtain a therapeutic effect. The oxidized glutathione and/or pharmaceutically acceptable salt and/or derivative is introduced along with an extender of their half life. The compds. of the invention may be used in the treatment of neoplasms, immune diseases, etc.

L36 ANSWER 30 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:350237 HCAPLUS

DOCUMENT NUMBER: 127:13137

TITLE: Retinoid induces growth inhibition of adult T-cell

leukemia cells

AUTHOR(S): Miyatake, Jun-Ichi; Maeda, Yasuhiro

CORPORATE SOURCE: Third Department of Internal Medicine, Kinki

University School of Medicine, Osakasayama, 589, Japan

SOURCE: Acta Medica Kinki University (1997), 22(1),

111-121

CODEN: AMKUDT; ISSN: 0386-6092

PUBLISHER: Kinki University Medical Association

DOCUMENT TYPE: Journal LANGUAGE: English

The effects of retinoic acid (RA) on the cell growth and expression of interleukin-2 (IL-2) receptor (IL-2 $R\alpha/p55$, Tac, CD25) by the human T lymphotropic virus type I pos. (HTLV-I(+)) T cell lines, HUT102 and ATL-2, were investigated. Incubation of these cells with RA resulted in marked growth inhibition and down-regulation of CD25 expression. Four clones of HUT102 cell lines were established by limiting dilution, and RA was shown to inhibit the growth and CD25 expression in three of these clones, but in the fourth. However, RA did not induce growth inhibition of the HTLV-I-neg. T cell lines, MOLT-4 and Jurkat, and of normal lymphocytes that had been stimulated with phytohemagglutinin. We hypothesized that the sensitivity to retinoids depends on an imbalance in intracellular redox potential. To examine the effect of exogenous thiol compds. for the growth inhibition of HTLV-I(+) T cell lines induced by RA, these cell lines were cultured with several thiol compds. (ATL-derived factor, thioredoxin, L-cystine and glutathione (GSH)), following the addition of RA in thiol -free medium. Unexpectedly, thiol compds. alone, when added after RA, did not restore the growth inhibition of HTLV-I(+) T cell lines induced by RA. However, when those cells were preincubated with thiol compds. for 24 h, no RA-induced growth inhibition was observed These findings suggest that intracellular reductive environments induced by thiol compds. are associated with resistance to RA of HTLV-I(+) T cells, and that thiol compds. may play an important role in HTLV-I(+) T cell proliferation.

REFERENCE COUNT: 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 31 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:153170 HCAPLUS

DOCUMENT NUMBER: 126:223448

TITLE: Decreased activity of inducible nitric oxide synthase

type 2 and modulation of the expression of glutathione S-transferase α , bcl-2, and

metallothioneins during the differentiation of CaCo-2

cells

AUTHOR(S): Vecchini, Françoise; Pringault, Eric; Billiar, Timothy

R.; Geller, David A.; Hausel, Pierrette; Felley-Bosco,

Emanuela

CORPORATE SOURCE: Institut de Pharmacologie et Toxicologie, Lausanne,

1005, Switz.

SOURCE: Cell Growth & Differentiation (1997), 8(2),

261-268

CODEN: CGDIE7; ISSN: 1044-9523

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

AB Reactive oxygen species modulate the cell growth of a wide variety of

mammalian cells. To determine whether oxidative metabolism is altered during the

differentiation process, we studied the expression of pro- and antioxidant proteins in proliferating and differentiated CaCo-2 cells, a human colon adenocarcinoma cell line. Nitric oxide synthase type 2 (iNOS) produces nitric oxide (NO). Depending on its rate of synthesis, NO may either promote cellular and DNA damage or reduce the ability of other free radicals to induce cell injury. Using Western and Northern blot anal. and arginine conversion assay, we demonstrate that the expression of iNOS decreases when cells undergo differentiation. This biol. event entails a diminished production of NO metabolites and correlates with the loss of activation of soluble guanylate cyclase activity. In differentiated cells, a 2-fold down-regulation of the nuclear factor kB activity was observed, suggesting that nuclear factor kB could be one of the iNOS gene regulatory factors in the CaCo-2 model. In parallel, we studied the expression of other antioxidant proteins including glutathione S-transferase α (GST α), bcl-2, and the metallothioneins (MTs). We show that the protein levels of $GST\alpha$ and MT increase during the differentiation of CaCo-2 cells, whereas bcl-2 levels decrease. Our investigation indicates that the expression of iNOS, GSTa, bcl-2, and MT is associated with the enterocytic differentiation. The shift in the expression of specific antioxidant genes during CaCo-2 cell differentiation may occur to avoid alterations in the cell redox potential.

L36 ANSWER 32 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:21606 HCAPLUS

DOCUMENT NUMBER: 126:152441

TITLE: Induction of p21 mediated by reactive oxygen species

formed during the metabolism of

aziridinylbenzoquinones by HCT116 cells

AUTHOR(S): Qiu, Xiaobo; Forman, Henry Jay; Schoenthal, Axel H.;

Cadenas, Enrique

CORPORATE SOURCE: Sch. Pharm., Univ. Southern California, Los Angeles,

CA, 90033, USA

SOURCE: Journal of Biological Chemistry (1996),

271(50), 31915-31921

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER: American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE: Journal LANGUAGE: English

AB Aziridinylbenzoquinones are a group of antitumor agents that elicit cytotoxicity by generating either alkylating intermediates or reactive oxygen species. The mechanism of toxicity may not always,

however, involve profound damage of cellular constituents, but may involve

a cytostatic effect through interference with the cell cycle. In this context, the authors have examined the induction of the cell cycle inhibitor p21 (WAF1, CIP1, or sdil), whose overexpression suppresses the growth of various tumor cells, in human tumor cells metabolizing 3,6-diaziridinyl-1,4-benzoquinone (DZQ) and its C2,C5-substituted derivs.: 2,5-bis-(carboethoxyamino) (AZQ) and 2,5-bis-(2-hydroxyethylamino) (BZQ). Both DZQ and AZQ were effectively activated by HCT116 human colonic carcinoma cells; the activation of the former involved largely a dicoumarol-sensitive activity, whereas that of the latter appeared to be accomplished primarily by one-electron transfer reductases. BZQ was not a substrate for the dicoumarol-sensitive enzyme in HCT116 cells. Cellular activation of the first two quinones was associated with formation of oxygen-centered radicals as detected by EPR in conjunction with the spin trap 5,5'-dimethyl-1-pyrroline-N-oxide. The redox transitions of DZQ involved hydroxyl radical formation and were strongly inhibited by catalase, whereas those of AZQ showed a strong superoxide anion component sensitive to superoxide dismutase. These signals were suppressed by N-acetylcysteine with concomitant production of a thiyl radical adduct. This suggests an effective electron transfer between the thiol and free radicals formed during the activation of these quinones. DZQ and AZQ induced significantly the expression of p21 in HCT116 cells, but a 10-fold higher concentration of AZQ was required to achieve the level of induction elicited by DZQ. BZQ had little effect on p21 expression. P21 induction at both mRNA and protein levels correlated with the inhibition of either cyclin-dependent kinase activity or cell proliferation. P21 induction elicited by the above quinones was inhibited by N-acetylcysteine, whereas the non-sulfur analog, N-acetylalanine, was without effect. Catalase and superoxide dismutase did not effect p21 induction by aziridinylbenzoquinones in HCT116 cells, thus suggesting that extracellular sources of oxygen radicals generated by plasma membrane reductases have no influence in the expression of this gene. Hydrogen peroxide, a product of quinone redox cycling, elicited an increase of p21 mRNA levels in HCT116 and K562 human chronic myelogenous leukemia cells. The latter lacks p53, one of the activators of p21 transcription, thus suggesting that p21 expression can be accomplished in a p53-independent manner in these cells. This study suggests that p21 induction is mediated by an increase in the cellular steady-state concentration of oxygen radicals and that the greater effectiveness in p21 induction by DZQ may be related to its efficient metabolism by NAD(P)H:quinone oxidoreductase activity in HCT116 cells. REFERENCE COUNT: 52 THERE ARE 52 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L36 ANSWER 33 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:936 HCAPLUS

DOCUMENT NUMBER: 126:84934

TITLE: Melatonin and oncostatic signal transduction: evidence

for a novel mechanism involving glutathione

and nitric oxide

AUTHOR(S): Blask, David E.; Wilson, Sean T.

CORPORATE SOURCE: Mary Imogene Bassett Hosp., Res. Inst., Cooperstown,

NY, 13326-1394, USA

SOURCE: Advances in Pineal Research (1994), 8,

465-471

CODEN: APIREW; ISSN: 0269-0071

PUBLISHER: Libbey
DOCUMENT TYPE: Journal
LANGUAGE: English

AB Melatonin is a unique neurohormone that has many diverse functions in

addition to its well-known role in biol. timekeeping. One of these functions is its ability to inhibit the promotion of tumor growth in vivo and in vitro, particularly of breast cancer cells. However, virtually nothing is known with respect to potential signal transduction pathways that may mediate melatonin's oncostatic action at the cellular level. Since one mechanism of tumor promotion may involve a prooxidant state of cancer cells, the authors have examined the role of the intracellular redox state in melatonin's mechanism of action in MCF-7 human breast cancer cells in vitro. Specifically, the authors report new evidence for a novel signal transduction mechanism mediating melatonin's oncostatic action involving the endogenous antioxidant glutathione (GSH) and a free radical species and newly discovered intracellular messenger mol., nitric oxide (NO). When the synthesis of either GSH or NO is inhibited, melatonin no longer exerts its antiproliferative effect, suggesting that these mols. play a critical role in transmitting melatonin's oncostatic message within breast cancer cells.

L36 ANSWER 34 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:756018 HCAPLUS

DOCUMENT NUMBER: 126:87554

TITLE: Oxidative inactivation of thioredoxin as a

cellular growth factor and protection by a Cys73

→ Ser mutation

AUTHOR(S): Gasdaska, John R.; Kirkpatrick, D. Lynn; Montfort,

William; Kuperus, Miles; Hill, Simon R.; Berggren,

Margareta; Powis, Garth

CORPORATE SOURCE: Arizona Cancer Center, Univ. Arizona Health Services

Center, Tucson, AZ, 85724-5024, USA

SOURCE: Biochemical Pharmacology (1996), 52(11),

1741-1747

CODEN: BCPCA6; ISSN: 0006-2952

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal
LANGUAGE: English

Thioredoxin (Trx) is a widely distributed redox protein that regulates several intracellular redox-dependent processes and stimulates the proliferation of both normal and tumor cells. We have found that when stored in the absence of reducing agents, human recombinant Trx undergoes spontaneous oxidation, losing its ability to stimulate cell growth, but is still a substrate for NADPH-dependent reduction by human thioredoxin reductase. There is a slower spontaneous conversion of Trx to a homodimer that is not a substrate for reduction by thioredoxin reductase and that does not stimulate cell proliferation. Both conversions can be induced by chemical oxidants and are reversible by treatment with the thiol reducing agent dithiothreitol. SDS-PAGE suggests that Trx undergoes oxidation to monomeric form(s) preceding dimer formation. recently shown by X-ray crystallog. that Trx forms a dimer that is stabilized by an intermol. Cys73-Cys73 disulfide bond. A Cys73 \rightarrow Ser mutant Trx (C73S) was prepared to determine the role of Cys73 in oxidative stability and growth stimulation. C73S was as effective as Trx in stimulating cell growth and was a comparable substrate for thioredoxin reductase. C73S did not show spontaneous or oxidant-induced loss of activity and did not form a dimer. suggest that Trx can exist in monomeric forms, some of which are mediated by Cys73 that do not stimulate cell proliferation but can be reduced by thioredoxin reductase. Cys73 is also involved in formation of an enzymically inactive homodimer, which occurs on long term

storage or by chemical oxidation Thus, although clearly involved in protein inactivation, Cys73 is not necessary for the growth stimulating activity of Trx.

L36 ANSWER 35 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:608659 HCAPLUS

DOCUMENT NUMBER: 125:271948

TITLE: Selenite and selenate inhibit human lymphocyte growth

via different mechanisms

AUTHOR(S): Spyrou, Giannis; Bjoernstedt, Mikael; Skog, Sven;

Homgren, Arne

CORPORATE SOURCE: Department of Medical Biochemistry and Biophysics,

Karolinska Institutet, Stockholm, S-171 77, Swed.

SOURCE: Cancer Research (1996), 56(19), 4407-4412

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

Selenium compds. like selenite and selenate have strong inhibitory effects, particularly on mammalian tumor cell growth by unknown mechanisms. We found that the addition of sodium selenite and sodium selenate inhibited the growth of human 3B6 and BL41 lymphocytes. Selenite was more potent because 10 µM selenite produced a growth inhibitory effect similar to that of 250 μM selenate. The mechanism of action of selenite and selenate appears to be different. 3B6 and BL41 cells treated with selenite accumulated in the S-phase; however, selenate caused an accumulation of cells in G2. Selenite-mediated growth inhibition was irreversible, although the effects of selenate could be reversed. Selenite, in contrast to selenate, is efficiently reduced by the thioredoxin system (thioredoxin, thioredoxin reductase, and NADPH). At concns. required to observe a similar effect on cell growth, the activity of thioredoxin reductase, recently shown to be a selenoprotein, increased in selenite-treated cells and decreased in selenate-treated cells. Ribonucleotide reductase activity was inhibited in an in vitro assay by selenite and selenodiglutathione but not by selenate. These results show that

selenite and selenate use different mechanisms to inhibit cell growth.

L36 ANSWER 36 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:263007 HCAPLUS

DOCUMENT NUMBER: 124:314095

TITLE: Reduction-oxidation (redox) state regulation

of extracellular matrix metalloproteinases and tissue

inhibitors in cardiac normal and transformed

fibroblast cells

AUTHOR(S): Tyagi, Suresh C.; G. Suresh Kumar; Borders, Susan CORPORATE SOURCE: Dalton Cardiovascular Research Center, University

Missouri-Columbia, Columbia, MO, 65212, USA

SOURCE: Journal of Cellular Biochemistry (1996),

61(1), 139-51

CODEN: JCEBD5; ISSN: 0730-2312

PUBLISHER: Wiley-Liss DOCUMENT TYPE: Journal LANGUAGE: English

AB Latent matrix metalloproteinases (MMPs) in normal myocardium are activated in end-stage heart failure. In vitro oxidized glutathione (GSSG) activates myocardial MMPs which contains a cysteine residue. In vivo GSSG induce the collagen lysis and cardiac dilatation. To assess whether thiol and non-thiol reducing agents have

direct effect on the interstitial human heart fibroblast (HHF) proliferation and MMP expression, HHF and polyoma virus transformed fibroblast cells were cultured with or without the thiol-containing reduced (GSH) or oxidized (GSSG) glutathiones , pyrrolidine dithiocarbamate (PDTC) and N-acetylcysteilne (NAC), and non-thiol ascorbic acid. After 100 µg/mL (.apprx.0.3 mM) GSH or PDTC treatment the proliferative (synthetic) phenotype of transformed fibroblast cells was changed to quiescent (contractile) phenotype. Also, after GSH, PDTC, and ascorbic acid treatment the medium was then analyzed for MMP activity by zymog. The results indicate reduction in MMP expression in transformed fibroblast cells after GSH and PDTC treatments and no effect after ascorbic acid treatment. Based on reverse zymog., we observed the level of tissue inhibitor of metalloproteinase (TIMP) at a decreased level in transformed cells. The effect of the reducing agent at the gene transcription was measured by estimating mRNA (Northern blot anal.) of MMP and of TIMP in the cells that were cultured in medium in the presence and absence of GSH. These results indicate that GSH induces MMP-2 and MMP-1 expression in normal HHF and that GSH reduces MMP-2 and MMP-1 in transformed fibroblast cells. After the treatment, the TIMP-2 level was repressed in normal HHF and TIMP-2 level increased in transformed fibroblast cells. These events are dependent on the nuclear transcription factor activity on the collagenase promoter in normal HHF cells. On the other hand, in polyoma transform fibroblast cells these events are not dependent on this collagenase promoter. These results suggest that oxidative environment induces normal HHF cell proliferation, and the reducing agent decreases normal HHF cell proliferation by inducing MMP and repressing TIMP gene transcription. In transformed cells reducing agents inhibit MMP expression and increase TIMP levels, which suggests a role of antioxidants in preventing tumorigenesis.

L36 ANSWER 37 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

1996:205604 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 124:313432

TITLE: Redox changes in normal and

neoplastic cells during the cell cycle. I.

Bioreduction of nitroxides by CHO cells with different

mitotic activity

AUTHOR (S):

Panz, Tadeusz

CORPORATE SOURCE:

Institute Molecular Biology, Jagiellonian University,

Krakow, Pol.

SOURCE:

Current Topics in Biophysics (1994), 18(2),

112-16

CODEN: CTOBEU; ISSN: 1232-9630

PUBLISHER:

Wydawnictwo Protext

Journal

DOCUMENT TYPE: LANGUAGE: English

Chinese Hamster Ovary (CHO) cells cultured in vitro and isolated in logarithmic phase of growth reduced spin probes at a lower rates than cells isolated during plateau phase of growth. This phenomenon was observed for nitroxides located in cell membranes and those penetrating into the cells. Blocking of electron transport in mitochondria with inhibitors slowed down the bioredn., whereas uncoupling of mitochondrial phosphorylation increased the rate of this process.

L36 ANSWER 38 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:170826 HCAPLUS

DOCUMENT NUMBER: 124:220550

TITLE: Treatment for atherosclerosis and other cardiovascular and inflammatory diseases with dithiocarboxylates and

dithiocarbamates which block VCAM-1 expression

INVENTOR(S): Medford, Russell M.; Alexander, R. Wayne;

Parthasarathy, Sampath; Khan, Bobby V.; Offermann,

Margaret K.

PATENT ASSIGNEE(S): Emory University, USA SOURCE: PCT Int. Appl., 99 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

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OTHER SOURCE(S): MARPAT 124:220550

AB Dithiocarboxylates, including dithiocarbamates, block the induced expression of the endothelial cell surface adhesion mol. VCAM-1, and are therefore useful in the treatment of cardiovascular disease, including atherosclerosis, as well as noncardiovascular inflammatory diseases that are mediated by VCAM-1. Identification of oxidized and unoxidized polyunsatd. fatty acids as direct mediators of VCAM-1 expression is described.

L36 ANSWER 39 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1996:46443 HCAPLUS

DOCUMENT NUMBER:

124:169927

TITLE:

Identification of proteins that are abnormally regulated in differentiated cultured human

keratinocytes

AUTHOR(S):

Olsen, Eydfinnur; Rasmussen, Hanne Holm; Celis, Julio

Ε.

CORPORATE SOURCE: Dep. Medical Biochem., Aarhus Univ., Aarhus, Den.

SOURCE: Electrophoresis (1995), 16(12), 2241-8

CODEN: ELCTDN; ISSN: 0173-0835

PUBLISHER: VCH
DOCUMENT TYPE: Journal
LANGUAGE: English

Comparison of the protein expression patterns of proliferating AB normal primary human keratinocytes plated in serum-free medium (SFKM) supplemented with epidermal growth factor (EGF) and bovine pituitary extract (BPE), and similar cultures induced to differentiate by the addition of Dulbecco's modified Eagle medium (DMEM), containing 10% fetal calf serum (FCS) revealed several known and unknown polypeptides that are abnormally regulated in the differentiated cells. Upregulated proteins included keratins (keratin 6, 10/11, 14 and 16), members of the S100 protein family (psoriasin, MRP8, MRP14 and S100c), actin-binding proteins (gelsolin and tropomyosin 9220), annexins (annexins IV and VIII), hsp28, the fatty acid binding protein 5 (FABP5) the squamous cell carcinoma (SCC) antigen, members of the 14-3-3 family, involucrin, E-cadherin, cystatin A, desmoglein and integrins $\alpha 2$ and $\beta 1$, as well as several proteins of as yet unknown identity. The highest upregulated proteins correspond to psoriasin (124.0 times), MRP8 (42.4 times), MRP14 (14.9 times), tropomyosin 9220 (11.5 times), involucrin (11.1 times), and FABP5 (9.1 times). FABP5, hsp28, and tropomyosin 9220 were also highly upregulated in quiescent keratinocytes indicating that their increased levels in the differentiated cells may be due to loss of proliferative activity. Highly downregulated proteins included PAI-2, tropomyosins 9213, 9121 and 9122, keratin 5, calnexin, 14-3-3 beta and eta, nucleoside diphosphate kinase A, Rho GDIs, hsp60, hnRNPs H and C2, α -enolase, eIF-4D, thioredoxin, annexins III and V, moesin, nucleolar protein B23, GST π and PCNA/cyclin. Both the high expression of keratin 6 and 16, which are markers for an alternative pathway of keratinocyte differentiation, as well as the extremely high upregulation of some members of the S100 protein family indicate that the cells differentiated via an abnormal pathway.

L36 ANSWER 40 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:28756 HCAPLUS

DOCUMENT NUMBER: 124:78608

TITLE: The organization of the human GSTP1-1 gene promoter

and its response to retinoic acid and cellular

redox status

AUTHOR(S): Xia, Chulin; Hu, Jiangting; Ketterer, Brian; Taylor,

John B.

CORPORATE SOURCE: Department Biochemistry Molecular Biology, University

College London, London, W1P 6DB, UK

SOURCE: Biochemical Journal (1996), 313(1), 155-61

CODEN: BIJOAK; ISSN: 0264-6021

PUBLISHER: Portland Press

DOCUMENT TYPE: Journal LANGUAGE: English

AB High levels of expression of GSTP1-1 are associated with cell proliferation, embryogenesis and malignancy. Given the role of glutathione S-transferase (GST) in detoxication, it is possible that GSTP1-1 evolved specifically to protect proliferating cells and share regulatory mechanisms with other cellular genes which are involved in cell division and tumorigenesis. We have previously shown that the expression of GSTP1 is suppressed by retinoic acid (RA) in the presence of the retinoic acid receptor (RAR) as a result of decreased

transcription from its promoter. Through deletion anal., we show here that the RA-RAR-dependent repression is mediated by the region -73 to +8. Further mutation anal. of this region indicates that the DNA sequence required for RA-RAR-dependent repression co-localizes with a consensus activator protein-1 (AP1) site essential for the promoter activity. degree of repression correlates with the residual activity of the AP1 site. There are two adjacent G/C boxes. The one immediately downstream from the AP1 site is not essential for the promoter activity, but mutation of the second, further downstream, impairs the promoter. On the other hand, mutation of either of these two G/C boxes has little effect on RA-RAR suppression. We also show that the expression of GSTP1 is regulated by the redox status of the cell. Using the chloramphenicol acetyltransferase assay system, we have demonstrated that treatment with H2O2 induced transcription from the promoter and that this effect can be blocked by pre-incubation with Nacetylcysteine (NAC). It was shown that the induction by H2O2 is mediated by trans-acting factor NF-kB (nuclear factor κB), via a putative NF-κB site, 'GGGACCCTCC', located from -96 to -86. Co-transfection with an NF-κB (p65) expression construct increased the promoter activity, an effect which could be blocked by co-transfection with an IkB (MAD-3) expression construct. Deletion of the NF-kB site abolished the effect of both H2O2 and co-transfection of NF- κ B. Interestingly, NAC is also an inducer for GSTP1. The effect of NAC was shown to be mediated largely by the AP1 site, since mutation of this site abolished the induction by NAC.

L36 ANSWER 41 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:830335 HCAPLUS

DOCUMENT NUMBER: 123:336694

TITLE: Influence of redox status of lymphocytes and

monocytes on HIV transcription and replication AUTHOR(S): Gougerot-Pocidalo, Marie-Anne; Aillet, Fabienne;

Virelizier, Jean-Louis; Israel, Nicole

CORPORATE SOURCE: INSERM, Hopital Bichat, Paris, Fr.

SOURCE: Immunobiology (Stuttgart) (1995), 193(2-4),

204-9

CODEN: IMMND4; ISSN: 0171-2985

PUBLISHER: Fischer

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review with 27 refs. Studies with antioxidants (NAC or BHA) revealed that NF-kB is involved in HIV virus replication in monocyte cell lines. In vivo results suggest that one of the limitations of antioxidant therapy might be that tissue macrophages multiply the virus actively. Also, BHA or NAC concns. able to partially inhibit HIV replication in latently infected mononuclear cells are deleterious to the immune system, as shown by the inhibition of interleukin-2-induced proliferation of human mononuclear cells and of inhibition of tumor necrosis factor secretion by monocytic cells.

L36 ANSWER 42 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:707270 HCAPLUS

DOCUMENT NUMBER: 123:195274

TITLE: Relationship between antioxidant systems,

intracellular thiols and DNA ploidy in liver of rats during experimental cirrhogenesis

AUTHOR(S): Sanz, Nuria; Diez-Fernandez, Carmen; Fernandez-Simon,

Lourdes; Alvarez, Alberto; Cascales, Maria

CORPORATE SOURCE: Facultad Farmacia, Univ. Complutense, Madrid, 28040,

Spain

SOURCE: Carcinogenesis (1995), 16(7), 1585-93

CODEN: CRNGDP; ISSN: 0143-3334

PUBLISHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

Hyperplastic nodular cirrhosis was induced in rats by long-term (6 mo) AB i.p. administration of thioacetamide at doses of 2.66 mmol/kg body weight, three times per wk. The survival rate of animals at the end of the treatment was 90%. To follow the temporal changes samples at 0, 7, 15, 30, 45, 60, 90, 150 and 180 days from rats during thioacetamide intoxication and from chronol. controls were obtained. The cirrhogenic ability of this treatment was assessed on the basis of morphol. changes: the development of macronodular cirrhosis and the appearance of fibrous septa of collagen through portal spaces. Parameters of liver injury and cholestasis were obtained by assaying the serum activities of isocitrate dehydrogenase and γ -glutamyltransferase. Enzymes and metabolites related to glutathione redox systems, as well as other antioxidant enzymes, were tested. Catalase and glutathione peroxidase, the two enzymes involved in the elimination of peroxides, and glutathione reductase decreased significantly at the end of the 6 mo of intoxication, while Cu-Zn and Mn superoxide dismutases increased progressively during the long-term thioacetamide treatment. Protein thiol levels profile showed a biphasic change increasing from the 7th day and were insensitive to the 30% depletion of intracellular glutathione (GSH). To study the relation of the intracellular thiols on the mechanisms of cell proliferation and differentiation during the cirrhogenic process, DNA content was assayed by flow cytometry in isolated hepatocytes, and DNA ploidy and distribution between G0-G1, S and G2 + M phases were determined Remarkable changes in relation to a sharp increase in diploid population from 7 to 180 days $(24.5\% \rightarrow 85.5\%)$, a pronounced decrease in polyploid populations (tetraploid + octoploid) in the same period $(73.7\% \rightarrow 12.3\%)$, and elevations in the populations in S phase (S1 + S2) were observed in thioacetamide-treated rats. The results obtained indicate that hepatocytes isolated from thioacetamide-treated rats showed a marked tendency to diploidy, an enhancement in DNA replication parallel to the hepatic content of protein sulfhydryl groups and a significant decline in antioxidant enzyme activities. The increase in protein thiols was independent of GSH level and of the thiol redox state.

L36 ANSWER 43 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1995:647796 HCAPLUS

TITLE: Increased levels of oxidized glutathione in

CD4+ lymphocytes associated with disturbed

intracellular redox balance in human
immunodeficiency virus type 1 infection

AUTHOR(S): Aukrust, Paal; Svardal, Asbjorn M.; Mueller, Fredrik;

Lunden, Bodil; Berge, Rolf K.; Ueland, Per M.;

Froeland, Stig S.

CORPORATE SOURCE: Clinical Immunology Infectious Diseases, Univ. Oslo,

Oslo, Norway

SOURCE: Blood (1995), 86(1), 258-61

CODEN: BLOOAW; ISSN: 0006-4971

PUBLISHER: Saunders
DOCUMENT TYPE: Journal
LANGUAGE: English

AB We investigated the intracellular glutathione redox status in isolated lymphocyte subpopulations and monocytes in patients with human immunodeficiency virus type 1 (HIV-1) infection and healthy controls. CD4+ lymphocytes from HIV-1-infected patients were primarily characterized by a substantial increase in oxidized glutathione levels and a considerable decrease in the ratio of reduced to total qlutathione, in most cases below 0.5 in patients with symptomatic HIV-1 infection, rather than decreased levels of reduced glutathione. The increase in oxidized glutathione was strongly correlated with low nos. of CD4+ lymphocytes in peripheral blood and impaired stimulated interleukin-2 production and proliferation in peripheral blood mononuclear cells, which is compatible with an immunopathogenic role for these redox disturbances. The HIV-1-infected patients with the most advanced clin. and immunol. disease were also characterized by an increase in levels of reduced glutathione in monocytes, suggesting that the glutathione redox cycle may be differentially regulated in CD4+ lymphocytes and monocytes. We could not confirm previous reports suggesting cysteine deficiency as a major cause of disturbed glutathione homeostasis during HIV-1 infection. The demonstrated glutathione abnormalities were correlated with raised serum levels of tumor necrosis factor α . These findings suggest that a therapeutical approach, which can restore the glutathione redox dysbalance in CD4+ lymphocytes and decrease the inflammatory stress, may be worthwhile exploring in HIV-1 infection.

L36 ANSWER 44 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1995:559396 HCAPLUS

DOCUMENT NUMBER:

122:312694

TITLE:

A CD4+ T cell line-secreted factor, growth promoting

for normal and leukemic B cells, identified as

thioredoxin

AUTHOR(S):

Rosen, Anders; Lundman, Pia; Carlsson, Mats; Bhavani,

Kasibhatla; Srinivasa, Bachally R.; Kjellstroem,

Gunilla; Nilsson, Kenneth; Holmgren, Arne

CORPORATE SOURCE:

Dep. Cell Biol., Univ. Linkoeping, Linkoeping, S-581

85, Swed.

SOURCE:

International Immunology (1995), 7(4),

625-33

CODEN: INIMEN; ISSN: 0953-8178

PUBLISHER:
DOCUMENT TYPE:

Oxford University Press Journal

LANGUAGE:

English

In this study, a B cell growth stimulatory factor, constitutively secreted by a human CD4+ T cell hybridoma clone, MP6, has been purified and characterized. Serum-free 24 h culture media from MP6 cells were collected, concentrated by ultrafiltration and separated by gel chromatog. Fractions were analyzed for stimulatory activity using [3H]thymidine incorporation in normal and leukemic (B-CLL) B cells as target cells. Activity was present in a 12 kDa protein peak. Upon storage this lost activity indicating that the factor was sensitive to air oxidation, a well-known property of mammalian thioredoxins (Trxs). Treatment of the inactive fraction with dithiothreitol restored full activity. When culture medium was analyzed with a RIA for human placenta Trx, the MP6 clone was shown to release 30-50 ng/mL per million cells during 24 h. The B cell stimulatory activity of the MP6 medium was removed by Sepharose-bound anti-human placenta Trx IgG and activity was recovered by elution from the antibodies. Furthermore, MP6 medium showed Trx activity with NADPH and Trx reductase using an insulin disulfide reduction assay. Starting from 5 L of serum-free MP6 conditioned medium, Trx was purified .apprx.100,000-fold. After gel electrophoresis banding, the material was analyzed by peptide sequencing and a full length sequence of an 104 amino acid long protein was obtained. This Trx sequence was identical to the previously published sequence of human Trx from HTLV-I transformed T cells, adult T cell leukemia-derived factor/Trx. A minor fraction (.apprx.30%) of the purified Trx showed alternative amino acids at eight positions; the relevance of which is discussed. MP6-derived Trx showed prominent growth stimulatory activity, measured as [3H] thymidine incorporation, and synergy was detected, particularly with IL-2, but also with IL-1 β , IL-4, IL-6, tumor necrosis factor- α , IFN- γ , low mol. weight B cell growth factor and antiCD40 mAbs. Interestingly, the promoter for the trx gene was recently reported to contain several sequence, motifs compatible with regulated inducible transcription, especially by cytokines. Together with the authors' previous results showing a cytokine/mitogen inducible autocrine secretion of Trx from B cells, the findings point to a crucial role for extracellular Trx in redox-controlled mechanisms of the B lymphocyte activation cascade.

L36 ANSWER 45 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1995:533909 HCAPLUS

DOCUMENT NUMBER:

122:283909

TITLE:

Analysis of studies related to tumorigenicity

induced by hydroquinone

AUTHOR (S):

Whysner, J.; Verna, L.; English, J. C.; Williams, G.

CORPORATE SOURCE:

Division Pathology and Toxicology, American Health

Foundation, Valhalla, NY, 10595, USA

SOURCE:

Regulatory Toxicology and Pharmacology (1995

), 21(1), 158-76

CODEN: RTOPDW; ISSN: 0273-2300

DOCUMENT TYPE:

Journal; General Review

LANGUAGE:

English

A review and discussion with many refs. which summarizes the bioassay data and analyzes information related to possible mechanisms for the tumorigenicity of hydroquinone (HQ). HQ produced renal adenomas in male F344 rats, and these tumors appeared to arise from areas of spontaneous progressive nephropathy; the nephropathy itself has been found to be enhanced by HQ. Other neoplasms were not confirmed to be causally related to HQ among the reported bioassays. In the male F344 rat, HQ administered alone was not DNA-reactive. HQ produced enhanced proliferation of renal tubular epithelium, presumably through toxicity involving glutathione conjugate formation. In the kidney, bone marrow, and other tissues, HQ may induce toxicity by redox cycling and lipid peroxidn. In bone marrow, HQ may produce microtubulin dysfunction, which is a plausible explanation for pos. cytogenetic tests, the only consistently pos. genotoxicity effect reported for HQ. Although HQ is a metabolic product of benzene, several lines of evidence suggest that the effects of HQ exposure are significantly different from those of benzene. Based upon the plausible mechanisms by which HQ may produce kidney tumors in male rats, the authors have concluded that occupational exposure levels of HQ are not predicted to be a cancer risk for humans.

L36 ANSWER 46 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1994:603247 HCAPLUS

DOCUMENT NUMBER:

121:203247

TITLE:

Influence of redox status of lymphocytes and

monocytes on HIV expression and immune functions. Evaluation in vitro of antioxidant molecules as

potential anti-HIV therapy

AUTHOR(S): Israel, N.; Gougerot-Pocidalo, M. A.; Aillet, F.;

Virelizier, J. L.

CORPORATE SOURCE: Unite d'Immunologie Virale, Inst. Pasteur, Paris,

75724, Fr.

SOURCE: Oxid. Stress, Cell Act. Viral Infect. (1994)

, 301-10 CODEN: 60KKAM

DOCUMENT TYPE: Conference LANGUAGE: English

The authors used BHA, a phenolic, lipid-soluble, chain-breaking antioxidant to show that peroxyl radical scavenging blocks NF-kB activation and HIV1 enhancer activity in PMA- or TNF-stimulated lymphoblastoid T (J.Jhan) and monocytic (U937) cells lines. The anti-oxidative effect of BHA was accompanied by an increase in thiol, but not glutathione , content in stimulated and unstimulated T cells, whereas TNF stimulation itself barely modified the cellular thiol level. Oxidative stress obtained by the addition of H2O2 to the culture medium of J.Jhan or U937 cells could not by itself induce NF-κB activation. These observations suggest that TNF and PMA do not lead to NF-κB activation through induction of changes in the cell redox status. Rather, TNF and PMA can exert a full activation effect only if cells are in a basal redox equilibrium tending towards oxidation since prior modification towards reduction by BHA treatment prevents their activation effects. The effects of BHA or NAC, a known glutathione precursor, were investigated also on the regulation of HIV1 expression in latently infected U1 cells and in the productively and chronically infected U937 cells. Both antioxidants inhibited TNF- or PMA-induced NF-kB activity in U1 cells in parallel with a partial decrease in induction of HIV replication. Both prevented the sustained NF- κB activity permanently induced by the virus in HIV chronically infected U937 but intriguingly did not modify HIV replication. This may be a limitation to potential antiviral effects of antioxidant therapies. Another limitation may be that antiviral (at least partially) concns. of NAC or BHA inhibited IL2-induced human PBMC proliferation and also secretion of TNF in PMA-stimulated U937 cells.

L36 ANSWER 47 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:602520 HCAPLUS

DOCUMENT NUMBER: 121:202520

TITLE: Abnormal redox regulation in HIV infection

and other immunodeficiency diseases

AUTHOR(S): Droge, W.; Eck, H. P.; Mihm, S.; Galter, D.

CORPORATE SOURCE: Div. Immunochem., Deutsches Krebsforschungszentrum,

Heidelberg, D-6900, Germany

SOURCE: Oxid. Stress, Cell Act. Viral Infect. (1994)

, 285-99 CODEN: 60KKAM

DOCUMENT TYPE: Conference; General Review

LANGUAGE: English

AB A review with 66 refs. HIV-infected persons at all stages of the disease have on the average markedly decreased plasma cystine and cysteine concns., decreased intracellular glutathione and elevated plasma glutamate levels. Elevated extracellular glutamate levels aggravate the cysteine deficiency since glutamate inhibits competitively the membrane transport of cystine. Lymphocyte functions in vitro are augmented even by moderate elevations of extracellular cysteine and inhibited by elevation

of the extracellular glutamate concns. A significant correlation between individual CD4+ T cel $\bar{1}$ nos. and individual cystine and glutamate levels has also been found in a cohort of HIV-infected persons, in healthy human blood donors, and in chimpanzees. CD8+ T cells showed no significant correlation. A rapid and significant decrease of plasma cysteine levels and increase of plasma glutamate was also found in rhesus macaques 2 wk after infection with the closely related SIVmac, but not in HIV-infected chimpanzees or SIVagm-infected African green monkeys. (The latter two species do not develop AIDS-like symptoms.). Elevated plasma glutamate levels were found to be neg. correlated with lymphocyte functions also in cancer patients. In view of the decreased levels of the bona fide antioxidants cysteine and qlutathione one may expect to find manifestations of oxidative damage. Indeed, elevated levels of malondialdehyde have been demonstrated, but the contribution of oxidative damage to the immunopathol. of HIV infection remains to be determined A cysteine deficiency is also expected to compromise certain glutathione-dependent immunol. functions, such as IL-2 dependent proliferation and activation of cytotoxic T cells. The activation of the transcription factor NFκB which controls the inducible transcription of several immunol. relevant genes, in contrast, was found to be neg. correlated with the extracellular cysteine supply. This indicates that the overactivation of several immunol. functions in the early stages of the disease, including the overexpression of an interleukin-2 receptor α -chain cleavage product, TNF α and β 2-microglobulin may also be the consequence of the HIV-induced cysteine deficiency. The replication of HIV-1, i.e. another gene under control of NFkB binding sites, was shown to be inhibited by cysteine or N-acetyl-cysteine (NAC). In view of the established cysteine and glutathione deficiency in HIV-infected persons, the authors have proposed to consider N-acetyl-cysteine for the treatment of these patients. NAC is a well established and safe drug with well documented toxicol. and pharmacokinetics.

L36 ANSWER 48 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:577403 HCAPLUS

DOCUMENT NUMBER: 121:177403

TITLE: Antioxidant and staurosporine inhibit stimulation of

the transcription regulator NF- κB following tumor necrosis factor treatment of chronic

B-leukemia cells

AUTHOR(S): Jabbar, Shireen A. B.; Hoffbrand, A. Victor;

Wickremasinghe, R. Gitendra

CORPORATE SOURCE: Department of Haematology, Royal Free Hospital School

of Medicine, London, NW3 2QG, UK

SOURCE: Leukemia Research (1994), 18(7), 523-30

CODEN: LEREDD; ISSN: 0145-2126

DOCUMENT TYPE: Journal LANGUAGE: English

AB B-chronic lymphocytic leukemia (B-CLL) and hairy cell leukemia cells (HCL) are refractory to stimulation by several cytokines which activate normal B-cells. However, tumor necrosis factor (TNF) promotes the proliferation of these cells. TNF regulates some of its cellular responses via the transcription factor NF- κ B. Using an electrophoretic mobility shift assay, we demonstrate that TNF treatment of B-CLL and HCL cells in vitro resulted in the augmentation of NF- κ B levels. In hemopoietic cell lines, TNF induction of NF- κ B is mediated via the generation of reactive oxygen intermediates and by the activation of protein kinase C (PKC). We have used activators and inhibitors of these pathways to unravel TNF signalling in the cells of ten

patients with B-CLL and two with HCL, using the increase in NF-kB levels following TNF treatment as an end point. Raising glutathione levels with N-acetyl cysteine substantially reduced NF-kB induction by TNF in two of four samples, as did treatment of cells with the antioxidant butylated-hydroxytoluene in all three samples These data suggest that redox mechanisms are involved in TNF signalling in these cells. Treatment with the PKC activator phorbol myristate acetate failed to activate NF-xB suggesting that this enzyme does not mediate the induction of NF-kB in these cells. However, the protein kinase inhibitor staurosporine inhibited TNF induction of NF-kB in four of five samples, suggesting that staurosporine-sensitive protein kinases (other than PKC) are involved in the signalling pathway. Our results suggest that PKC-independent pathways, including pathways sensitive to redox reagents, mediate the induction of NF-κB by TNF in chronic B-leukemia cells. Addnl., these data suggest that defects in PKC-mediated pathways may contribute to the general reluctance of B-CLL and HCL cells to respond to mitogenic signals.

L36 ANSWER 49 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:505379 HCAPLUS

DOCUMENT NUMBER: 121:105379

TITLE: Oxidative damage and repair in the developing nervous

system

AUTHOR(S): Verity, M. Anthony

CORPORATE SOURCE: Division Neuropathology and Brain Research Institute,

UCLA Center the Health Sciences, Los Angeles, CA,

90024-1732, USA

SOURCE: Neurotoxicology (1994), 15(1), 81-91

CODEN: NRTXDN; ISSN: 0161-813X

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review with 88 refs. Excessive production of reactive oxygen species (ROS) is a recognized cause of cell injury. In contrast to such well recognized cell injury, oxidative stress plays a role in cell proliferation , differentiation and tumor promotion. This review examines the role of oxidative stress in initiating and promoting the establishment of normal or abnormal neuronal patterns and subsequent neurogenesis within the central and peripheral nervous system. In particular, the role of apoptosis in both normal and abnormal neuronal development and maturation will be examined with special reference to the induction of apoptotic cell death

following abusive ligand-induced ion movements. The interaction of oxidant stress and immediate-early response gene activation is discussed with further reference to the induction of apoptosis. While glutamate receptor activation appears mandatory for coordinate maturation and neuritogenesis, such neuronal survival and differentiation is intimately dependent upon the intracellular glutathione redox potential, maintained by cystine uptake. Selected examples of reactive oxygen species induced injury pertaining to developmental neurotoxicol. are presented and include starvation, irradiation injury and glutamate excitotoxicity.

L36 ANSWER 50 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:429020 HCAPLUS

DOCUMENT NUMBER: 121:29020

TITLE: Pathobiological effects of acetaldehyde in cultured

human epithelial cells and fibroblasts

AUTHOR(S): Grafstroem, Roland C.; Dypbukt, Jeannette, M.;

Sundqvist, Kristina; Atzori, Luigi; Nielsen, Inge;

Curren, Rodger D.; Harris, Curtis C.

CORPORATE SOURCE: Inst. Environ. Med., Karolinska Inst., Stockholm,

S-171 77, Swed.

SOURCE: Carcinogenesis (1994), 15(5), 985-90

CODEN: CRNGDP; ISSN: 0143-3334

DOCUMENT TYPE: Journal LANGUAGE: English

AB The ability of acetaldehyde, a respiratory carcinogen present in tobacco smoke and automotive emissions, to affect cell viability, thiol status and intracellular Ca2+ levels and to cause DNA damage and mutations has been studied using cultured human cells. Within a concentration range of 3-100 mM, a 1 h exposure to acetaldehyde decreases colony

survival and inhibits uptake of the vital dye neutral red in bronchial epithelial cells. Acetaldehyde also causes both DNA interstrand cross-links and DNA protein cross-links whereas no DNA single strand breaks are detected. The cellular content of glutathione is also decreased by acetaldehyde, albeit, without concomitant changes in the glutathione redox status or in the content of protein Transient or sustained increases in cytosolic Ca2+ occur within minutes following exposure of cells to acetaldehyde. Moreover, acetaldehyde significantly decreases the activity of the DNA repair enzyme O6-methylguanine-DNA methyltransferase. Finally, a 5 h exposure to acetaldehyde causes significant levels of 6-thioguanine resistance mutations in an established mutagenesis model involving skin fibroblasts. The results indicate that mM concns. of acetaldehyde cause a wide range of cytopathic effects associated with multistep carcinogenesis. fact that acetaldehyde, in relation to its cytotoxicity, causes comparatively higher genotoxicity and inhibits DNA repair more readily than other major aldehydes in tobacco smoke and automotive emissions is discussed.

L36 ANSWER 51 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1993:515121 HCAPLUS

DOCUMENT NUMBER: 119:115121

TITLE: Augmentation of cytostatic effect of recombinant human

lymphotoxin and involvement of glutathione

redox cycle

AUTHOR(S): Matsunaga, K.; Mashiba, H.

CORPORATE SOURCE: Div. Immunol., Natl. Kyushu Cancer Cent., Fukuoka,

812, Japan

SOURCE: European Cytokine Network (1992), 3(3),

307-11

CODEN: ECYNEJ; ISSN: 1148-5493

DOCUMENT TYPE: Journal LANGUAGE: English

The effect of buthionine sulfoximine (BSO), an inhibitor of glutathione biosynthesis, in combined use with a nitrosourea derivative, ACNU, on the cytostatic effect of recombinant human lymphotoxin (rhLT) was studied in vitro. The simultaneous addition of 0.02 mM or 0.5 mM BSO and rhLT slightly augmented the inhibition of Meth A tumor cell proliferation. A similar tendency was observed when the target cells were treated with 0.02 mM or 0.5 mM BSO for 24 h prior to the addition of rhLT. A marked augmentation of the antiproliferative effect was obtained when the target cells were treated in vitro with 0.005 mM or 0.02 mM BSO prior to the addition of 0.02 mM or 0.1 mM BSO and rhLT. The addition of ACNU simultaneously with rhLT to BSO-treated cells also augmented the antiproliferative effect.

These results suggest that the **glutathione redox** cycle is closely related to the mechanism of LT-induced cytotoxicity.

L36 ANSWER 52 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1992:39448 HCAPLUS

DOCUMENT NUMBER: 116:39448

TITLE: Red cell regulation of tumor necrosis

factor-induced human neutrophil cytostatic activity

AUTHOR(S): Shau, Hungyi

CORPORATE SOURCE: Sch. Med., UCLA, Los Angeles, CA, 90024-1782, USA

SOURCE: Cancer Communications (1991), 3(9), 283-6

CODEN: CNCMET; ISSN: 0955-3541

DOCUMENT TYPE: Journal LANGUAGE: English

AB **Tumor** necrosis factor (TNF) activates polymorphonuclear neutrophils (PMN) to suppress **tumor** cell **proliferation**

. This cytostatic activity could be blocked by the addition of red blood cells (RBC) into the assay. TNF-induced PMN cytostatic activity was mediated by hydrogen peroxide. RBC have two major pathways to detoxify H2O2, one by catalase and the other by the glutathione redox cycle. Therefore, the catalase inhibitor 3-amino-1,2,4-triazole (AT) and the glutathione inhibitor N-ethylmaleimide (NE) were used to assess the role of each anti-oxidant in protecting the tumor target cells. RBC, depleted of catalase by AT, no longer protected Raji tumor cells from PMN cytostatic activity. However, depletion of reduced glutathione by NE had no effect on RBC protection of tumor target cells. Thus, RBC

TNF-activated PMN, and the protection is a function of catalase, but not glutathione.

L36 ANSWER 53 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1991:441503 HCAPLUS

DOCUMENT NUMBER: 115:41503

TITLE: Augmented antiproliferative effect in

combined use of human lymphotoxin with a nitrosourea

derivative, ACNU, and the involvement of

glutathione redox cycle

can protect tumor cells from cytostatic activity mediated by

AUTHOR(S): Mashiba, Harukazu; Matsunaga, Keiko; Kakutani, Tetsu CORPORATE SOURCE: Div. Immunol., Natl. Kyushu Cancer Cent., Fukuoka,

815, Japan

SOURCE: International Journal of Immunopharmacology (

1991), 13(4), 333-8

CODEN: IJIMDS; ISSN: 0192-0561

DOCUMENT TYPE: Journal LANGUAGE: English

AB The cytotoxic or cytostatic effect of the combined use of human lymphotoxin (LT) with 1-(4-amino-2-methyl-5-pyrimidinyl)methyl-3-(2-chloroethyl)-3-nitrosourea-HCl (ACNU) on L cells or Meth A tumor cells was studied. Simultaneous addition of LT derived from a human lymphoid cell line with ACNU (200 or 500 μg/mL) significantly augmented the cytotoxic effect. Similar augmented inhibition was obtained when LT was added to ACNU-treated L cells. The pretreatment of Meth A tumor cells with ACNU (25 or 50 μg/mL) augmented recombinant human LT-mediated cytostasis. However, the addition of glutathione (1.0 mg/mL) to ACNU-treated Meth A tumor cells significantly nullified the augmented antiproliferative effect of LT (10 U/mL). These results suggest that augmentation of the antiproliferative effect on tumor cells could be induced

through the combined use of LT with ACNU by lowering the intracellular level of glutathione.

L36 ANSWER 54 OF 54 HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1988:400358 HCAPLUS

DOCUMENT NUMBER: 109:358

TITLE: Toxic effects of acute glutathione depletion

by buthionine sulfoximine and

dimethylfumarate on murine mammary carcinoma

cells

AUTHOR(S): Dethlefsen, L. A.; Lehman, C. M.; Biaglow, J. E.;

Peck, V. M.

CORPORATE SOURCE: Health Sci. Cent., Univ. Utah Health, Salt Lake City,

UT, 84132, USA

SOURCE: Radiation Research (1988), 114(2), 215-24

CODEN: RAREAE; ISSN: 0033-7587

DOCUMENT TYPE: Journal LANGUAGE: English

Glutathione depletion to .simeq.5% of control for 48 h or longer by 0.05 mM L-buthionine sulfoximine (BSO) led to appreciable toxicity for the 66 murine mammary carcinoma cells growing in vitro [L. A. Dethlefsen et al., 1986]. Such toxicity in normal, proliferating cells in vivo would be undesirable. Thus the toxic effects after acute GSH depletion to .simeq.5% of control by BSO plus dimethylfumarate (DMF) were evaluated in these same 66 cells to determine if this anti-proliferative effect could be minimized. Two hours of 0.025 mM DMF reduced GSH to 45% of control, while 6 h of 0.05 mM BSO reduced it to 16%. However, BSO (6 h) plus DMF (2 h) and BSO (24 h) plus DMF (2 h) reduced GSH to 4 and 2%, resp. The incorporation (15-min pulses) of radioactive precursors into protein and RNA were unaffected by these treatment protocols. In contrast, cell growth was only modestly affected, but the incorporation of [3H] thymidine into DNA was reduced to 64% of control by the BSO (24 h) plus DMF (2 h) protocol even though it was unaffected by the BSO (6 h) plus DMF (2 h) treatment. However, the aerobic radiation response, as measured by cell survival, was not modified at doses of either 4.0 or 8.0 Gy. The growth rates of treated cultures, after drug removal, quickly returned to control rates and the resynthesis of GSH in cells from both protocols was also rapid. The GSH levels after either protocol were slightly above control by 12 h after drug removal, dramatically over control (.simeq.200%) by 24 h, and back to normal by 48 Thus even a relatively short treatment with BSO and DMF resulting in a GSH depletion to 2-5% of control had a marked effect on DNA synthesis and plating efficiency and a modest effect on cellular growth. Presumably the antiproliferative effects are due to a depletion of nuclear GSH with the subsequent inhibition of the GSH/glutaredoxin-mediated conversion of ribonucleotides to deoxyribonucleotides. However, even after extended treatment, upon drug removal, GSH was rapidly resynthesized and cellular DNA synthesis and growth quickly resumed.

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                    ?NEOPLAS? OR ?TUMOR? OR ?TUMOUR?)
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48 SEA FILE=HCAPLUS ABB=ON L30 AND ?THIOL?
54 SEA FILE=HCAPLUS ABB=ON L30 AND (PRD<19990216 OR PD<19990216)
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L35
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L37
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L38
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=> d ibib abs 138 1-67

L38 ANSWER 1 OF 67 MEDLINE on STN DUPLICATE 1 ACCESSION NUMBER: 2005266362 MEDLINE

ACCESSION NUMBER: 2005266362 MEDLI DOCUMENT NUMBER: PubMed ID: 15792952

TITLE: Mechanistic studies on a novel, highly potent

gold-phosphole inhibitor of human glutathione

reductase.

AUTHOR: Deponte Marcel; Urig Sabine; Arscott L David; Fritz-Wolf

Karin; Reau Regis; Herold-Mende Christel; Koncarevic Sasa; Meyer Markus; Davioud-Charvet Elisabeth; Ballou David P;

Williams Charles H Jr; Becker Katja

CORPORATE SOURCE: Interdisciplinary Research Center, Justus Liebig

University, D-35392 Giessen, Germany.

CONTRACT NUMBER: GM11106 (NIGMS)

SOURCE: Journal of biological chemistry, (2005 May 27) 280 (21)

20628-37. Electronic Publication: 2005-03-24.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200508

ENTRY DATE:

Entered STN: 20050524

Last Updated on STN: 20050816

Entered Medline: 20050815

The homodimeric flavoprotein qlutathione reductase (GR) is a AB central player of cellular redox metabolism, connecting NADPH to

the large pool of redox-active thiols. In this work, the inhibition of human GR by a novel gold-phosphole inhibitor (GoPI) has been studied in vitro. Two modes of inhibition are observed, reversible inhibition that is competitive with GSSG followed by irreversible inhibition. When approximately 1 nm GoPI is incubated with NADPH-reduced GR (1.4 nm) the enzyme becomes 50% inhibited. This appears to be the most potent stable inhibitor of human GR to date. Analyzing the monophasic oxidative half-reaction of reduced GR with GSSG at pH 6.9 revealed a K(d)((app)) for GSSG of 63 microm, and a k((obs)max) of 106 s(-1) at 4 degrees C. The reversible inhibition by the gold-phosphole complex [{1-phenyl-2,5-di(2-pyridyl)phosphole}AuCl] involves formation of a complex at the GSSG-binding site of GR (K(d) = 0.46 microm) followed by nucleophilic attack of an active site cysteine residue that leads to covalent modification and complete inactivation of the enzyme. Data from titration spectra, molecular modeling, stopped-flow, and steady-state kinetics support this theory. In addition, covalent binding of the inhibitor to human GR was demonstrated by mass spectrometry. The extraordinary properties of the compound and its derivatives might be exploited for cell biological studies or medical applications, e.g. as an anti-tumor or antiparasitic drug. Preliminary experiments with glioblastoma cells cultured in vitro indicate an anti-

proliferative effect of the inhibitor in the lower micromolar range.

L38 ANSWER 2 OF 67

MEDLINE on STN

DUPLICATE 2

ACCESSION NUMBER: DOCUMENT NUMBER:

IN-PROCESS 2005250952 PubMed ID: 15890017

TITLE:

Differential susceptibility of nonmalignant human breast

epithelial cells and breast cancer cells to

thiol antioxidant-induced G(1)-delay.

AUTHOR:

Menon Sarita G; Coleman Mitchell C; Walsh Susan A; Spitz

CORPORATE SOURCE:

Douglas R; Goswami Prabhat C

Free Radical and Radiation Biology Program, Department of Radiation Oncology, University of Iowa, Iowa City, IA

52242, USA.

CONTRACT NUMBER:

CA66081 (NCI)

HL51469 (NHLBI)

SOURCE:

Antioxidants & redox signalling, (2005 May-Jun) 7 (5-6)

711-8.

Journal code: 100888899. ISSN: 1523-0864.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

NONMEDLINE; IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE: Entered STN: 20050514

Last Updated on STN: 20050607

AB Reactive oxygen species (ROS) and ROS signaling have been implicated in a variety of human pathophysiological conditions that involve aberrant cellular proliferation, particularly cancer. We

hypothesize that intracellular redox state differentially affects cell-cycle progression in nonmalignant versus malignant cells. The thiol antioxidant, N-acetyl-L-cysteine (NAC), was used to alter intracellular redox state in nonmalignant human breast epithelial (MCF-10A) and breast cancer cells (MCF-7 and MDA-MB-231). Treatment of cells with NAC resulted in significant augmentation of intracellular small-molecular-weight thiols, glutathione and cysteine. In addition, NAC treatment decreased oxidation of a prooxidant-sensitive dye in MCF-10A cells, but not in MDA-MB-231 and MCF-7 cells. NAC -induced shifts in intracellular redox state toward a more reducing environment caused G(1) delays in MCF-10A cells without causing any significant changes in MCF-7 and MDA-MB-231 cell-cycle progression. NAC treatment of MCF-10A (but not MCF-7 and MDA-MB-231) was accompanied by a decrease in cyclin D1 and an increase in p27 protein levels, which correlated with increased retinoblastoma protein hypophosphorylation. These results show differential redox control of progression from G(1) to S in nonmalignant versus malignant cells and support the hypothesis that loss of a redox control of the cell cycle could contribute to aberrant proliferation seen in cancer cells.

L38 ANSWER 3 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2005382451 EMBASE

TITLE: Proceedings from the "Third International Conference on

Mechanism of Action of Nutraceuticals".

AUTHOR: Mandel S.; Packer L.; Youdim M.B.H.; Weinreb O.

CORPORATE SOURCE: O. Weinreb, Department of Pharmacology, Technion-Faculty of

Medicine, P.O.B. 9697, Haifa 31096, Israel.

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SOURCE: Journal of Nutritional Biochemistry, (2005) Vol. 16, No. 9,

pp. 513-520.

Refs: 56

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COUNTRY: United States

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 017 Public Health, Social Medicine and Epidemiology

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20050915

Last Updated on STN: 20050915

AB The "Third International Conference on Mechanisms of Action of Nutraceuticals" (ICMAN 3) was held to bring investigators from around the world together to find answers and share experience relevant to the role of nutraceuticals in health and disease. Dietary supplements are currently receiving recognition as being beneficial in coronary heart disease, cancer, osteoporosis and other chronic and degenerative diseases such as diabetes, Parkinson's and Alzheimer's diseases. This gave impetus to investigate the mechanisms of action of nutraceuticals and related bioactive compounds in disease pathologies. Many lines of evidence indicate that the mechanistic actions of natural compounds involve a wide array of biological processes, including activation of antioxidant defenses, signal transduction pathways, cell survival-associated gene expression, cell proliferation and differentiation and preservation of mitochondrial integrity. Furthermore,

many of these compounds exert anti-inflammatory actions through inhibition of oxidative stress-induced transcription factors (e.g., NF- κ B, AP-1), cytotoxic cytokines and cyclooxygenase-2. It appears that these properties play a crucial role in the protection against the pathologies of numerous age-related or chronic diseases. This review summarizes the latest research finding in functional foods and micronutrients in the promotion of health and reduction of risk for major chronic diseases as presented in this symposium. .COPYRGT. 2005 Elsevier Inc. All rights reserved.

L38 ANSWER 4 OF 67 MEDLINE on STN

ACCESSION NUMBER: 2005312756 MEDLINE DOCUMENT NUMBER: PubMed ID: 15746213

TITLE: Extracellular cysteine/cystine redox regulates

the p44/p42 MAPK pathway by metalloproteinase-dependent

epidermal growth factor receptor signaling.

AUTHOR: Nkabyo Yvonne S; Go Young-Mi; Ziegler Thomas R; Jones Dean

Graduate Program in Molecular and Systems Pharmacology,

Emory University, Atlanta, Georgia 30322, USA.

CONTRACT NUMBER: DK-55850 (NIDDK)

ES-011195 (NIEHS)

CORPORATE SOURCE:

SOURCE: American journal of physiology. Gastrointestinal and liver

physiology, (2005 Jul) 289 (1) G70-8. Electronic

Publication: 2005-03-03.

Journal code: 100901227. ISSN: 0193-1857.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200508

ENTRY DATE: Entered STN: 20050618

Last Updated on STN: 20050803 Entered Medline: 20050802

AB Previous research shows that stimulation of proliferation of colon carcinoma (Caco-2) cells by a more reduced extracellular cysteine/cystine (Cys/CySS) redox state occurs with no apparent effect on intracellular glutathione and that this stimulation is lost on addition of epidermal growth factor. The purpose of the present study was to determine whether a more reduced extracellular Cys/CySS redox state activates the mitogenic p44/p42 mitogen-activated protein kinase (MAPK) pathway and whether this is signaled through the epidermal growth factor receptor (EGFR). Caco-2 cells were exposed to a range of physiological extracellular redox conditions from -150 to 0 mV. In the absence of added growth factors, the most reduced (-150 mV) redox state induced an 80% increase in EGFR phosphorylation, and this was followed by a marked increase in phosphorylation of p44/p42 MAPK. Inhibitors of EGFR (AG1478) and p44/p42 MAPK (U0126) phosphorylation blocked redox-dependent p44/p42 phosphorylation, indicating that signaling occurred by EGFR. These effects were inhibited by pretreatment with a nonpermeant alkylating agent, showing that signaling involved thiols accessible to the extracellular space. The EGFR ligand TGF-alpha was increased in culture medium at more reduced redox states. Redox-dependent phosphorylation of EGFR was completely prevented by a metalloproteinase inhibitor (GM6001), and an antibody to TGF-alpha partially inhibited the phosphorylation of p44/p42 MAPK by redox. Thus the data show that a redox -dependent activation of metalloproteinase can stimulate the mitogenic p44/p42 MAPK pathway by a TGF-alpha-dependent mechanism. Because Cys

availability and Cys/CySS redox are dependent on nutrition, disease, and environmental exposures, the results suggest that cell proliferation could be influenced physiologically by Cys-dependent redox effects on growth factor signaling pathways.

L38 ANSWER 5 OF 67 MEDLINE ON STN
ACCESSION NUMBER: 2005222593 MEDLINE
DOCUMENT NUMBER: PubMed ID: 15684606

TITLE: The thioredoxin reductase/thioredoxin

system: novel redox targets for cancer

therapy.

AUTHOR: Biaglow John E; Miller Richard A

CORPORATE SOURCE: Department of Radiation Oncology and Biochemistry,

University of Pennsylvania Medical School, Philadelphia

19104, USA.. Biaglow@xrt.upenn.edu

SOURCE: Cancer biology & therapy, (2005 Jan) 4 (1) 6-13.

Electronic Publication: 2004-01-08. Ref: 79 Journal code: 101137842. ISSN: 1538-4047.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200509

ENTRY DATE: Entered STN: 20050429

Last Updated on STN: 20050916 Entered Medline: 20050915

AB Thioredoxin reductase (TRX) is a selenoprotein that reduces oxidized protein substrates in an NADPH-dependent process (cf. Fig. 1). The thioredoxins (TX) are a family of small redox active proteins that undergo reversible oxidation/reduction and help to maintain the redox state of cells. TX serves as a cofactor in many TRX-catalyzed reductions in a manner similar to qlutathione (GSH) in thioltransferase reactions. For example, TX is a cofactor in protein disulfide reduction and DNA synthesis, but independently, it inhibits apoptosis, stimulates cell proliferation and angiogenesis, and increases transcription factor activity. The role of the TRX/TX system is limited by its reducing capacity as well as the additional supply of electrons in the form of NADPH provided by hexose monophosphate shunt (HMPS). TX is limited by the reduction capacity of its vicinal sulfhydryls and needs a source of electrons from the HMPS and TRX- coupled system to reduce disulfides. Oxidized TX is reduced by TRX and NADPH. Several lines of evidence suggest that the coupled HMPS/TRX/TX system represents an important target for cancer therapy. TX overexpression has been reported in several malignancies and may be associated with aggressive tumor growth and poor survival. In some cells, TX is an important factor in conferring resistance to chemotherapy and in stimulating production of hypoxia-inducible factor (HIF-1). Several inhibitors of the TRX/TX system have been evaluated in experimental cancer models: these include HMPS inhibitors, carbohydrate analogues, NADP synthesis blockers, vicinal thiol reactants, cisplatin, and TRX inhibitors. More recently, the targeted anti-cancer agent motexafin gadolinium has been identified. Motexafin gadolinium is a redox mediator that selectively localizes to cancer cells, and reacts with reducing metabolites and vicinal thiols to generate reactive oxygen species that ultimately block the TRX enzyme as well as the analogous glutaredoxin activity. In cell and animal models, motexafin gadolinium is directly cytotoxic to various tumor cells and

enhances the activity of radiation therapy and chemotherapy. This drug is now in a broad range of clinical trials investigating its therapeutic potential when used as a single agent or in combination with either chemotherapy or radiation therapy. Promising clinical activity has been reported in a clinical trial with motexafin gadolinium and whole brain radiation therapy for treatment of brain metastases from solid tumors. These findings suggest that the TRX/TX system may represent an attractive target for development of new cancer therapeutics.

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on STN

ACCESSION NUMBER: 2005306014 EMBASE

TITLE: Extracellular cysteine/cystine redox regulates

the p44/p42 MAPK pathway by metalloproteinase-dependent

epidermal growth factor receptor signaling.

AUTHOR: Nkabyo Y.S.; Go Y.-M.; Ziegler T.R.; Jones D.P.

CORPORATE SOURCE: D.P. Jones, Dept. of Medicine, Whitehead Biomedical

Research Center, Emory Univ., 615 Michael St., Atlanta, GA

30322, United States. dpjones@emory.edu

SOURCE: American Journal of Physiology - Gastrointestinal and Liver

Physiology, (2005) Vol. 289, No. 1 52-1, pp. G70-G78.

Refs: 42

ISSN: 0193-1857 CODEN: APGPDF

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

037 Drug Literature Index

048 Gastroenterology

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20050721

Last Updated on STN: 20050721

Previous research shows that stimulation of proliferation of AB colon carcinoma (Caco-2) cells by a more reduced extracellular cysteine/cystine (Cys/CySS) redox state occurs with no apparent effect on intracellular glutathione and that this stimulation is lost on addition of epidermal growth factor. The purpose of the present study was to determine whether a more reduced extracellular Cys/CySS redox state activates the mitogenic p44/p42 mitogen-activated protein kinase (MAPK) pathway and whether this is signaled through the epidermal growth factor receptor (EGFR). Caco-2 cells were exposed to a range of physiological extracellular redox conditions from -150 to 0 mV. In the absence of added growth factors, the most reduced (-150 mV) redox state induced an 80% increase in EGFR phosphorylation, and this was followed by a marked increase in phosphorylation of p44/p42 Inhibitors of EGFR (AG1478) and p44/p42 MAPK (U0126) phosphorylation blocked redox-dependent p44/p42 phosphorylation, indicating that signaling occurred by EGFR. These effects were inhibited by pretreatment with a nonpermeant alkylating agent, showing that signaling involved thiols accessible to the extracellular space. The EGFR ligand $TGF-\alpha$ was increased in culture medium at more reduced redox states. Redox-dependent phosphorylation of EGFR was completely prevented by a metalloproteinase inhibitor (GM6001), and an antibody to $TGF-\alpha$ partially inhibited the phosphorylation of p44/p42 MAPK by redox. Thus the data show that a redox-dependent activation of metalloproteinase can stimulate the mitogenic p44/p42 MAPK pathway by a TGF- α -dependent mechanism. Because Cys availability and Cys/CySS redox are

dependent on nutrition, disease, and environmental exposures, the results suggest that cell **proliferation** could be influenced physiologically by Cys-dependent **redox** effects on growth factor signaling pathways. Copyright .COPYRGT. 2005 the American Physiological Society.

L38 ANSWER 7 OF 67 MEDLINE ON STN
ACCESSION NUMBER: 2004109223 MEDLINE
DOCUMENT NUMBER: PubMed ID: 14998722

TITLE: Thiol antioxidant and thiol-reducing

agents attenuate 15-deoxy-delta 12,14-prostaglandin

J2-induced heme oxygenase-1 expression.

AUTHOR: Liu Jean-Dean; Tsai Shu-Huei; Lin Shyr-Yi; Ho Yuan-Soon;

Hung Ling-Fang; Pan Shiann; Ho Feng-Ming; Lin Chun-Mao;

Liang Yu-Chih

CORPORATE SOURCE: College of Medicine, Taipei Medical University, Taipei,

Taiwan.. ycliang@tmu.edu.tw

SOURCE: Life sciences, (2004 Mar 26) 74 (19) 2451-63.

Journal code: 0375521. ISSN: 0024-3205.

PUB. COUNTRY: England: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200404

ENTRY DATE: Entered STN: 20040305

Last Updated on STN: 20040403 Entered Medline: 20040402

AB Heme oxygenase-1 (HO-1) is induced as a beneficial and adaptive response in cells and tissues exposed to oxidative stress. Herein we examined how various eicosanoids affect the induction of HO-1, and the possible mechanism underlying 15-deoxy-Delta(12,14) - prostaglandin J(2) (15d-PGJ(2))-induced HO-1 expression. PGH(2), PGD(2) and its metabolites of the PGJ(2) series, and PGA(1) markedly induced the protein expression of HO-1. Arachidonic acid (AA), docosahexaenoic acid (DHA), PGE(2), PGF(2 alpha), and thromboxane B(2) (TXB(2)) were shown to have no effect on the induction of HO-1. 15d-PGJ(2) was the most potent activator achieving significance at 5 microM. Although 15d-PGJ(2) significantly activated the MAPKs of JNK and ERK, the activation of JNK and ERK did not contribute to the induction of HO-1 as determined using transfection of dominant-negative plasmids and MAPKs inhibitors. Additional experiment indicated that 15d-PGJ(2) induced HO-1 expression through peroxisome proliferator-activated receptor (PPAR)-independent pathway. 15d-PGJ(2) significantly decreased the intracellular level of reduced glutathione; and the thiol antioxidant, N-acetyl-L-cysteine (NAC), and the thiol-reducing agent, dithiothreitol (DTT), inhibited the induction of HO-1 by 15d-PGJ(2). Finally, NAC and DTT exhibited significant inhibition of HO-1 mRNA and HO-1 promoter reporter activity induced by 15d-PGJ(2). These results suggest that thiol antioxidant and reducing agents attenuate the expression of HO-1 induced by 15d-PGJ(2), and that the cellular thiol-disulfide redox status may be linked to HO-1 activation.

L38 ANSWER 8 OF 67 MEDLINE on STN DUPLICATE 3

ACCESSION NUMBER: 2005028939 MEDLINE DOCUMENT NUMBER: PubMed ID: 15480664

TITLE: Sodium selenite induces apoptosis in acute promyelocytic

leukemia-derived NB4 cells by a caspase-3-dependent mechanism and a **redox** pathway different from that

of arsenic trioxide.

AUTHOR: Zuo Lu; Li Jian; Yang Yang; Wang Xuan; Shen Ti; Xu Cai-min;

Zhang Zhi-nan

CORPORATE SOURCE: Department of Hematology, Peking Union Medical College

Hospital, Peking Union Medical College and Chinese Academy of Medical Sciences, 100730, Beijing, People's Republic of

China.

SOURCE: Annals of hematology, (2004 Dec) 83 (12) 751-8. Electronic

Publication: 2004-10-06.

Journal code: 9107334. ISSN: 0939-5555.

PUB. COUNTRY: Germany: Germany, Federal Republic of
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200503

ENTRY DATE: Entered STN: 20050120

Last Updated on STN: 20050318 Entered Medline: 20050317

Two relatively recent discoveries stand behind our current effort to AB investigate the effects of the chemopreventive agent, selenium, on the proliferation and survival of NB4 cells. The first is that certain selenium compounds such as sodium selenite have pro-oxidant ability to catalyze the oxidation of thiols and simultaneously generate superoxide. The second lies in the exquisite susceptibility of NB4 cells to arsenic trioxide-induced, reactive oxygen species (ROS) -mediated apoptosis due to less efficiency of the cellular defense system. In this study, we demonstrated that sodium selenite could induce apoptosis in NB4 cells via the classic mitochondrial pathway involving caspase-3 activation and Bcl-2 cleavage. An increase in the basal cellular glutathione (GSH) content rendered NB4 cells resistant to arsenic trioxide, but could sensitize NB4 cells to sodium selenite. Moreover, combined treatment of NB4 cells with all- trans retinoic acid (ATRA) at low concentration and sodium selenite exhibited a synergistic effect on apoptosis induction. Together, our results suggest that selenite is a promising candidate for treatment of acute promyelocytic leukemia (APL) and the mechanism underlying its anticancer effects warrants further investigation.

L38 ANSWER 9 OF 67 MEDLINE on STN DUPLICATE 4

ACCESSION NUMBER: 2004141888 MEDLINE DOCUMENT NUMBER: PubMed ID: 14988435

TITLE: Glutathione metabolism and its implications for

health.

AUTHOR: Wu Guoyao; Fang Yun-Zhong; Yang Sheng; Lupton Joanne R;

Turner Nancy D

CORPORATE SOURCE: Faculty of Nutrition, Texas A&M University, College

Station, TX, 77843, USA.. g-wu@tamu.edu

CONTRACT NUMBER: P30-ES09106 (NIEHS)

R01CA61750 (NCI)

SOURCE: Journal of nutrition, (2004 Mar) 134 (3) 489-92. Ref: 31

Journal code: 0404243. ISSN: 0022-3166.

(Investigators: Lupton J R, TX A&M U, College Station)

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals; Space Life Sciences

ENTRY MONTH: 200404

ENTRY DATE:

Entered STN: 20040324

Last Updated on STN: 20040420 Entered Medline: 20040419

Glutathione (gamma-glutamyl-cysteinyl-glycine; GSH) is the most AB abundant low-molecular-weight thiol, and GSH/glutathione disulfide is the major redox couple in animal cells. The synthesis of GSH from glutamate, cysteine, and glycine is catalyzed sequentially by two cytosolic enzymes, gamma-glutamylcysteine synthetase and GSH synthetase. Compelling evidence shows that GSH synthesis is regulated primarily by gamma-glutamylcysteine synthetase activity, cysteine availability, and GSH feedback inhibition. Animal and human studies demonstrate that adequate protein nutrition is crucial for the maintenance of GSH homeostasis. In addition, enteral or parenteral cystine, methionine, N-acetyl-cysteine, and L-2-oxothiazolidine-4carboxylate are effective precursors of cysteine for tissue GSH synthesis. Glutathione plays important roles in antioxidant defense, nutrient metabolism, and regulation of cellular events (including gene expression, DNA and protein synthesis, cell proliferation and apoptosis, signal transduction, cytokine production and immune response, and protein glutathionylation). Glutathione deficiency contributes to oxidative stress, which plays a key role in aging and the pathogenesis of many diseases (including kwashiorkor, seizure, Alzheimer's disease, Parkinson's disease, liver disease, cystic fibrosis, sickle cell anemia, HIV, AIDS, cancer, heart attack, stroke, and diabetes). New knowledge of the nutritional regulation of GSH metabolism is critical for the development of effective strategies to improve health and to treat these diseases.

MEDLINE on STN L38 ANSWER 10 OF 67 ACCESSION NUMBER: 2003423750 MEDLINE DOCUMENT NUMBER: PubMed ID: 12796500

TITLE: A 16-residue peptide fragment of macrophage migration

> inhibitory factor, MIF-(50-65), exhibits redox activity and has MIF-like biological functions.

AUTHOR: Nquyen Mai Tuyet; Beck Jurgen; Lue Hongqi; Funfziq Helge;

Kleemann Robert; Koolwijk Pieter; Kapurniotu Aphrodite;

Bernhagen Jurgen

CORPORATE SOURCE: Division of Biochemistry and Molecular Cell Biology,

Institute of Biochemistry, University Hospital RWTH Aachen,

D-52074 Aachen, Germany.

SOURCE: Journal of biological chemistry, (2003 Sep 5) 278 (36)

33654-71. Electronic Publication: 2003-06-09.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200310

ENTRY DATE: Entered STN: 20030911

> Last Updated on STN: 20031008 Entered Medline: 20031007

AB Macrophage migration inhibitory factor (MIF) is a cytokine that participates in the host inflammatory response. A Cys-Xaa-Xaa-Cys (CXXC) -based thiol-protein oxidoreductase activity of MIF is associated with certain biological functions. Peptides spanning the CXXC region of thiol-protein oxidoreductases retain some biochemical properties of the full-length protein. We report on the characterization of CXXC-spanning MIF-(50-65) and its serine variant, C57S/C60S-MIF-(50-65). Following disulfide-mediated cyclization, MIF-(50-65) adapted a

beta-turn conformation comparable with that of beta-turn-containing cyclo-57,60-[Asp57,Dap60]MIF-(50-65). MIF-(50-65) had a redox potential E'0 of -0.258 V and formed mixed disulfides with glutathione and cysteine. MIF-(50-65) but not C57S/C60S-MIF-(50-65) had oxidoreductase activity in vitro. Intriquingly, MIF-(50-65) exhibited MIF-like cellular activities. The peptide but not its variant had glucocorticoid overriding and proliferation -enhancing activity and stimulated ERK1/2 phosphorylation. MIF-(50-65) and its variant bound to the MIF-binding protein JAB1 and enhanced cellular levels of p27Kip1. As the peptide and its variant were endocytosed at similar efficiency, sequence 50-65 appears sufficient for the JAB1-related effects of MIF, whereas other activities require CXXC. Cyclo-57,60-[Asp57,Dap60]MIF-(50-65) activated ERK1/2, indicating that CXXC-dependent disulfide and beta-turn formation is associated with an activity-inducing conformation. We conclude that CXXC and sequence 50-65 are critical for the activities of MIF. MIF-(50-65) is a surprisingly short sequence with MIF-like functions that could be an excellent molecular template for MIF therapeutics.

L38 ANSWER 11 OF 67 MEDLINE on STN DUPLICATE 5

ACCESSION NUMBER: 2003197407 MEDLINE DOCUMENT NUMBER: PubMed ID: 12574159

TITLE: Rapid induction of cell death by selenium-compromised

thioredoxin reductase 1 but not by the fully active

enzyme containing selenocysteine.

AUTHOR: Anestal Karin; Arner Elias S J

CORPORATE SOURCE: Medical Nobel Institute for Biochemistry, Department of

Medical Biochemistry and Biophysics, Karolinska Institute,

171 77 Stockholm, Sweden.

SOURCE: Journal of biological chemistry, (2003 May 2) 278 (18)

15966-72. Electronic Publication: 2003-02-06.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200306

ENTRY DATE: Entered STN: 20030429

Last Updated on STN: 20030618 Entered Medline: 20030617

AB Mammalian thioredoxin reductases are selenoproteins. For native catalytic activity, these enzymes utilize a C-terminal -Gly-Cys-Sec-Gly-COOH sequence (where Sec is selenocysteine) forming a redox active selenenylsulfide/selenolthiol motif. A range of cellular systems depend upon or are regulated by thioredoxin reductase and its major protein substrate thioredoxin, including apoptosis signal-regulating kinase 1, peroxiredoxins, methionine sulfoxide reductase, and several transcription factors. Cytosolic thioredoxin reductase 1 (TrxR1) is moreover inhibited by various electrophilic anticancer compounds. TrxR1 is hence generally considered to promote cell viability. However, several recent studies have suggested that TrxR1 may promote apoptosis, and the enzyme was identified as GRIM-12 (gene associated with retinoid interferon-induced mortality 12). Transient transfection with GRIM-12/TrxR1 was also shown to directly induce cell death. To further analyze such effects, we have here employed lipid-mediated delivery of recombinant TrxR1 preparations into human A549 cells, thereby bypassing selenoprotein translation to facilitate assessment of the protein-related effects on cell viability. We found that selenium-deficient TrxR1, having a two-amino acid-truncated C-terminal -Gly-Cys-COOH motif, rapidly induced cell death (38 +/- 29% apoptotic cells after 4 h; p < 0.005 compared with controls). Cell death induction was also promoted by selenium-compromised TrxRl derivatized with either cis-diamminedichloroplatinum (II) (cisplatin) or dinitrophenyl moieties but not by the structurally related non-selenoprotein **glutathione** reductase. In contrast, TrxRl with intact selenocysteine could not promote cell death. The direct cellular effects of selenium-compromised forms of TrxRl may be important for the pathophysiology of selenium deficiency as well as for the efficacy of **antiproliferative** drugs targeting the selenocysteine moiety of this enzyme.

L38 ANSWER 12 OF 67 MEDLINE on STN ACCESSION NUMBER: 2003119084 MEDLINE DOCUMENT NUMBER: PubMed ID: 12626594

TITLE: The cytokine macrophage migration inhibitory factor reduces

pro-oxidative stress-induced apoptosis.

AUTHOR: Nguyen Mai Tuyet; Lue Hongqi; Kleemann Robert; Thiele

Michael; Tolle Gabriele; Finkelmeier Doris; Wagner Eva;

Braun Andrea; Bernhagen Jurgen

CORPORATE SOURCE: Laboratory of Biochemistry, Institute for Interfacial

Engineering, University of Stuttgart and Fraunhofer Institut fur Grenzflachen-und Bioverfahrenstechnik,

Stuttgart, Germany.

SOURCE: Journal of immunology (Baltimore, Md.: 1950), (2003 Mar

15) 170 (6) 3337-47.

Journal code: 2985117R. ISSN: 0022-1767.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200306

ENTRY DATE: Entered STN: 20030314

Last Updated on STN: 20030626 Entered Medline: 20030625

AB The cytokine macrophage migration inhibitory factor (MIF) exhibits proand anti-inflammatory activities and regulates cell proliferation and survival. We investigated the effects of MIF on apoptosis. As MIF exhibits oxidoreductase activity and participates in regulating oxidative cell stress, we studied whether MIF could affect oxidative stress-induced apoptosis. We demonstrated that MIF exhibits antiapoptotic activity in various settings. MIF suppressed camptothecin-induced apoptosis in HeLa and Kym cells and HL-60 promyeloblasts. Both exogenous MIF and endogenous MIF, induced following overexpression through tetracycline (tet) gene induction, led to significant suppression of apoptosis. Apoptosis reduction by MIF was also observed in T cells. A role for MIF in redox stress-induced apoptosis was addressed by comparing the effects of rMIF with those of the oxidoreductase mutant C60SMIF. Endogenous overexpression of C60SMIF was similar to that of MIF, but C60SMIF did not suppress apoptosis. Exogenous rC60SMIF inhibited apoptosis. A role for MIF in oxidative stress-induced apoptosis was directly studied in HL-60 leukocytes and tet-regulated HeLa cells following thiol starvation or diamide treatment. MIF protected these cells from redox stress-induced apoptosis and enhanced cellular glutathione levels. As overexpressed C60SMIF did not protect tet-regulated HeLa cells from thiol starvation-induced apoptosis, it seems that the redox motif of MIF is important for this function. Finally, overexpression of MIF inhibited phosphorylation of endogenous c-Jun induced by thiol starvation, indicating that

MIF-based suppression of apoptosis is mediated through modulation of c-Jun N-terminal kinase activity. Our findings show that MIF has potent antiapoptotic activities and suggest that MIF is a modulator of pro-oxidative stress-induced apoptosis.

L38 ANSWER 13 OF 67 MEDLINE on STN DUPLICATE 6

ACCESSION NUMBER: 2002730644 MEDLINE DOCUMENT NUMBER: PubMed ID: 12494466

TITLE: Prostaglandin J2 metabolites inhibit aromatase activity by

redox-sensitive mechanisms: potential implications

for breast cancer therapy.

AUTHOR: Winnett Georgia; van Hagen Daphne; Schrey Michael

CORPORATE SOURCE: Endocrinology and Metabolic Medicine, Division of Medicine,

Faculty of Medicine, Imperial College of Science Technology

and Medicine, London, United Kingdom.

SOURCE: International journal of cancer. Journal international du

cancer, (2003 Feb 20) 103 (5) 600-5. Journal code: 0042124. ISSN: 0020-7136.

PUB. COUNTRY:

United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200302

ENTRY DATE: Entered STN: 20021221

Last Updated on STN: 20030207 Entered Medline: 20030206

AB The mechanisms by which prostaglandin (PG)J(2) metabolites inhibit tumorigenicity are poorly understood but may involve thiol reactivity or peroxisome proliferator-activated receptor (PPAR) -dependent pathways. Because aromatase is an important therapeutic target in breast cancer treatment, we have investigated the effect of PGJ(2) metabolites on aromatase activity and evaluated a potential role for redox status during PGJ(2) metabolite action. 15-deoxy-Delta(12,14)PGJ(2) (15d-PGJ(2)) and 9-deoxy-Delta(9,12)13,14dihydroPGD(2) (Delta(12)PGJ(2)) caused dose-dependent inhibition of both pre-induced aromatase activity in human breast fibroblasts and MDA MB 231 breast cancer cells and of constitutive aromatase activity in JEG-3 choriocarcinoma cells. Structure-activity studies showed that this inhibition was mimicked by 4-cyclopentene-1,3-dione but not by the PPARgamma agonist troglitazone nor the eicosanoids PGE(2) or arachidonic acid. The thiol oxidants diamide and H(2)O(2) simulated the inhibitory action of 15d-PGJ(2) on aromatase activity, whereas the glutathione (GSH) repletor and antioxidant N-acetyl-cysteine (NAC) reversed these actions of 15d-PGJ(2) and H(2)O(2) on aromatase. 15d-PGJ(2) also caused a direct dose-dependent inhibition of aromatase activity in JEG-3 cell sonicates, which was also reversed in the presence of GSH. Kinetic analysis of this 15d-PGJ(2)-induced inhibition of cell-free aromatase indicated the involvement of a non-competitive mechanism possibly resulting from direct thiol-targeted alkylation of the enzyme. These redox -sensitive, PPARgamma-independent actions of 15d-PGJ(2) on aromatase activity demonstrate a novel therapeutic potential for such cyclopentenone PGs in breast cancer treatment. Copyright 2002 Wiley-Liss, Inc.

L38 ANSWER 14 OF 67 MEDLINE on STN DUPLICATE 7

ACCESSION NUMBER: 2004006646 MEDLINE DOCUMENT NUMBER: PubMed ID: 14703721

TITLE: Activation of DNA biosynthesis in human hepatoblastoma

HEPG2 cells by the nitric oxide donor, sodium

nitroprusside.

AUTHOR: Sokolowska Maria; Rokita Hanna; Wlodek Lidia

CORPORATE SOURCE: Institute of Medical Biochemistry, Collegium Medicum,

Jagiellonian University, 31-034 Cracow, Poland.

SOURCE: Fundamental & clinical pharmacology, (2003 Oct) 17 (5)

599-607.

Journal code: 8710411. ISSN: 0767-3981.

PUB. COUNTRY: England: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200406

ENTRY DATE: Entered STN: 20040106

Last Updated on STN: 20040603 Entered Medline: 20040602

AB The role of nitric oxide (NO) in carcinogenesis is controversial as it has been shown to both stimulate and inhibit tumour growth. Also, there are contradictory opinions regarding the effects of NO on the proliferation of normal and tumour cells. The aim of our study was to use an in vitro model to determine the influence of exogenous NO donors on DNA biosynthesis by measuring [3H] thymidine incorporation in human hepatoblastoma cells (HepG2). studies were conducted with the following NO precursors: sodium nitroprusside (SNP), S-nitrosoglutathione, and nitroglycerine (NTG). Out of all three NO donors, SNP increased NO levels and strongly stimulated DNA biosynthesis. A SNP concentration of 150 microM induced optimal NO levels necessary for the activation of DNA biosynthesis. Lower levels of DNA biosynthesis (118% increase over the control) were observed in the presence of NTG, whereas S-nitrosoglutathione had no effect. Antioxidants such as thiol-containing drugs, N -acetylcysteine and tocopherol, proved to be the most efficient co-activators of SNP-induced DNA synthesis. On the other hand, supplementing the SNP-containing medium with compounds that induce oxidative stress and lower the level of -SH groups such as hydrogen peroxide, doxorubicin, and N-ethylmaleimide, led to the inhibition of DNA synthesis. Therefore, our results firmly confirm the hypothesis that biological effects of exogenous NO donors depends on the redox status of the cell.

L38 ANSWER 15 OF 67 MEDLINE ON STN ACCESSION NUMBER: 2003528606 MEDLINE DOCUMENT NUMBER: PubMed ID: 14606649

TITLE: Redox modulation of NF-kappaB nuclear

translocation and DNA binding in metastatic melanoma. The role of endogenous and gamma-glutamyl transferase-dependent

oxidative stress.

AUTHOR: Dominici Silvia; Visvikis Athanase; Pieri Lisa; Paolicchi

Aldo; Valentini Marta A; Comporti Mario; Pompella Alfonso

CORPORATE SOURCE: Department of Experimental Pathology, University of Pisa

Medical School, Pisa, Italy.

SOURCE: Tumori, (2003 Jul-Aug) 89 (4) 426-33.

Journal code: 0111356. ISSN: 0300-8916.

PUB. COUNTRY: Italy

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200312

ENTRY DATE: Entered STN: 20031111

Last Updated on STN: 20031219 Entered Medline: 20031202

AIMS AND BACKGROUND: The transcription factor NF-kappaB is implicated in AB the expression of genes involved in cell proliferation, apoptosis and metastasis. In melanoma, high constitutive levels of NF-kappaB activation are usually observed. NF-kappaB is regulated by oxidation/reduction (redox) processes, and the occurrence of constitutive oxidative stress in melanoma cells has been documented. Recent studies of our laboratories showed that the membrane-bound gamma-glutamyl transferase (GGT) enzyme activity--expressed by a number of malignancies, including melanoma -- can act as a basal source of superoxide, hydrogen peroxide and other prooxidants. METHODS: In the present study we utilized the 2/60 clone of Me665/2 human metastatic melanoma, which displays high levels of GGT activity, in order to verify if the presence of this enzyme--through the promotion of redox processes--may influence the activation status of NF-kappaB. The latter was evaluated by determining the nuclear translocation of the p65 subunit (by immunoblot), the DNA binding of NF-kappaB (by electrophoretic mobility shift assay) and its transcriptional activity (by gene transactivation studies). RESULTS: Me665/2/60 cells displayed a basal production of hydrogen peroxide. Stimulation of GGT activity by its substrates glutathione and glycyl-glycine caused additional production of hydrogen peroxide, up to levels approx. double the basal levels. Nuclear translocation of the NF-kappaB p65 subunit, DNA-binding and gene transactivation were thus investigated in Me665/2/60 cells whose GGT activity was modulated by means of substrates or inhibitors. Stimulation of GGT activity resulted in increased nuclear translocation of p65, while on the other hand NF-kappaB DNA binding and gene transactivation were paradoxically decreased. NF-kappaB DNA binding could be restored by treating cell lysates with the thiol-reducing agent dithiothreitol (DTT). Treatment of cells with exogenous hydrogen peroxide did not affect NF-kappaB activation status. CONCLUSIONS: Altogether, the data obtained indicate that GGT activity may impair the redox status of thiols that is critical for NF-kappaB DNA binding and gene transactivation, through the production of prooxidant species allegedly distinct from hydrogen peroxide. GGT activity therefore appears to be an additional factor in modulation of NF-kappaB transcriptional activity in melanoma, capable of hindering NF-kappaB DNA binding even in conditions where continuous oxidative stress would favor NF-kappaB nuclear translocation.

L38 ANSWER 16 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

ACCESSION NUMBER: 2003417596 EMBASE

TITLE: Redox state of glutathione and

thioredoxin in differentiation and apoptosis.

AUTHOR: Watson W.H.; Chen Y.; Jones D.P.

CORPORATE SOURCE: W.H. Watson, Department of Biochemistry, Emory University

School of Medicine, Atlanta, GA, United States

SOURCE: BioFactors, (2003) Vol. 17, No. 1-4, pp. 307-314.

Refs: 29

ISSN: 0951-6433 CODEN: BIFAEU

COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

> 029 Clinical Biochemistry

LANGUAGE: English

ENTRY DATE: Entered STN: 20031030

Last Updated on STN: 20031030

DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER

L38 ANSWER 17 OF 67 MEDLINE on STN DUPLICATE 8

ACCESSION NUMBER: 2003051328 MEDLINE PubMed ID: 12560100 DOCUMENT NUMBER:

TITLE: Redox catalysts as sensitisers towards oxidative

AUTHOR: Giles Niroshini M; Gutowski Nick J; Giles Gregory I; Jacob

Claus

School of Chemistry, University of Exeter, Stocker Road, CORPORATE SOURCE:

Exeter EX4 4QD, UK.

FEBS letters, (2003 Jan 30) 535 (1-3) 179-82. SOURCE:

Journal code: 0155157. ISSN: 0014-5793.

PUB. COUNTRY:

Netherlands

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200303

ENTRY DATE:

Entered STN: 20030202

Last Updated on STN: 20030322 Entered Medline: 20030321

The predominance of oxidative stress in many tumour cell AB environments provides a means to selectively target these cells via protein oxidation. The zinc fingers of transcription factors utilise cysteine thiols for structural zinc coordination. Redox control of DNA binding regulates transcription and therefore the overall rates of proliferation, apoptosis and necrosis in the carcinoma. We report here the adverse effects of glutathione peroxidase (GPx) mimics towards zinc finger motifs and PC12 cell survival. Nanomolar catalyst concentrations facilitated H202-induced oxidation of an Spl transcription factor fragment. In PC12 cells GPx catalysis triggered a significant increase in cell death, correlating with severity of oxidative stress. As a consequence, we conclude that GPx mimics are potential chemotherapeutic agents.

L38 ANSWER 18 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2003486063 EMBASE

TITLE: Glutamine and KGF each regulate extracellular thiol

> /disulfide redox and enhance proliferation in Caco-2 cells.

AUTHOR: Jonas C.R.; Gu L.H.; Nkabyo Y.S.; Mannery Y.O.; Avissar

N.E.; Sax H.C.; Jones D.P.; Ziegler T.R.

CORPORATE SOURCE: T.R. Ziegler, General Clinical Research Center, Emory Univ.

Hospital, 1364 Clifton Rd., Atlanta, GA 30322, United

States. tzieg01@emory.edu

SOURCE: American Journal of Physiology - Regulatory Integrative and

Comparative Physiology, (2003) Vol. 285, No. 6 54-6, pp.

R1421-R1429. Refs: 42

ISSN: 0363-6119 CODEN: AJPRDO

COUNTRY: United States DOCUMENT TYPE: Journal; Article FILE SEGMENT:

016 Cancer

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20040105

Last Updated on STN: 20040105

AΒ Glutamine (Gln) and keratinocyte growth factor (KGF) each stimulate

intestinal epithelial cell growth, but regulatory mechanisms are not well understood. We determined whether Gln and KGF alter intra- and extracellular thiol/disulfide redox pools in Caco-2 cells cultured in oxidizing or reducing cell medium and whether such redox variations are a determinant of proliferative responses to these agents. Cells were cultured over a physiological range of oxidizing to reducing extracellular thiol/disulfide redox (E(h)) conditions, obtained by varying cysteine (Cys) and cystine (CySS) concentrations in cell medium. Cell proliferation was determined by 5-bromo-2-deoxyuridine (BrdU) incorporation. Gln (10 mmol/l) or KGF (10 $\mu q/l)$ did not alter BrdU incorporation at reducing E(h) (-131 to -150 mV), but significantly increased incorporation at more oxidizing E (h) (Gln at 0 to - 109 mV; KGF at -46 to -80 mV). Cellular glutathione/glutathione disulfide (GSH/GSSG) E(h) was unaffected by Gln, KGF, or variations in extracellular Cys/CySS E(h). Control cells largely maintained extracellular E(h) at initial values after 24 h (-36 to -136 mV). However, extracellular E(h) shifted toward a narrow physiological range with Gln and KGF treatment (Gln -56 to -88 mV and KGF -76 to -92 mV, respectively; P < 0.05 vs. control). The results indicate that thiol/disulfide redox state in the extracellular milieu is an important determinant of Caco-2 cell proliferation induced by Gln and KGF, that this control is independent of intracellular GSH redox status, and that both Gln and KGF enhance the capability of Caco-2 cells to modulate extremes of extracellular redox.

L38 ANSWER 19 OF 67 MEDLINE on STN DUPLICATE 9

ACCESSION NUMBER: 2002409353 MEDLINE DOCUMENT NUMBER:

PubMed ID: 12163679

TITLE: Oxidation of the glutathione/glutathione

disulfide redox state is induced by cysteine deficiency in human colon carcinoma HT29 cells.

AUTHOR: Miller Lauren T; Watson Walter H; Kirlin Ward G; Ziegler

Thomas R; Jones Dean P

CORPORATE SOURCE: Department of Biochemistry, Emory University School of

Medicine, Atlanta, GA 30322, USA.

CONTRACT NUMBER: DK55850 (NIDDK)

> ES011195 (NIEHS) ES09047 (NIEHS)

SOURCE: Journal of nutrition, (2002 Aug) 132 (8) 2303-6.

Journal code: 0404243. ISSN: 0022-3166.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200209

ENTRY DATE: Entered STN: 20020807

> Last Updated on STN: 20030312 Entered Medline: 20020904

AB Glutathione (GSH) has a central role in the maintenance of the thiol-disulfide redox state in mammalian cells. GSH synthesis can be physiologically limited by the availability of cysteine (Cys), and Cys and its precursors are variable in the human diet. The purpose of this study was to determine the effect of severe Cys deficiency and readdition of Cys on the redox state of the GSH/ glutathione disulfide (GSSG) pool in human colon carcinoma HT29 cells. Cells were cultured in Cys- (and cystine-)limiting medium for 48 h followed by culture in medium containing either Cys or cystine for 24 h. GSH and GSSG were measured by HPLC. Cys limitation decreased cellular

GSH and GSSG concentrations with an associated >80 mV oxidation of the GSH/GSSG redox state. Upon addition of either Cys or its disulfide cystine (CySS), redox of GSH/GSSG recovered in 4 h, whereas GSH concentration continued to increase over 12 h. Maximal GSH concentrations attained were 200% of control cell values. These results show that severe Cys deficiency can have marked effects on cellular redox state but that redox recovers rapidly upon resupply. The magnitude of oxidation during Cys limitation in this cell model is sufficient to result in a >100-fold change in the reduced/oxidized ratio of redox-sensitive dithiol /disulfide motifs in proteins. If redox changes occur in vivo in association with variations in dietary Cys and its precursors, these changes could have important physiologic effects through altered redox signaling and control of cell proliferation and apoptosis.

L38 ANSWER 20 OF 67 MEDLINE on STN DUPLICATE 10

ACCESSION NUMBER: 2002687773 MEDLINE DOCUMENT NUMBER: PubMed ID: 12446207

TITLE: Extracellular thiol/disulfide redox

state affects proliferation rate in a human colon

carcinoma (Caco2) cell line.

AUTHOR: Jonas Carolyn R; Ziegler Thomas R; Gu Li H; Jones Dean P CORPORATE SOURCE: Department of Biochemistry, Emory University School of

Medicine, Atlanta, GA, USA.

CONTRACT NUMBER: M01 RR00039 (NCRR)

R01 ES011195 (NIEHS) R01 ES09047 (NIEHS) R01DK55850 (NIDDK) T32DK07732 (NIDDK)

SOURCE: Free radical biology & medicine, (2002 Dec 1) 33 (11)

1499-506.

Journal code: 8709159. ISSN: 0891-5849.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200305

ENTRY DATE: Entered STN: 20021214

Last Updated on STN: 20030529 Entered Medline: 20030528

Redox mechanisms function in regulation of cell growth, and AB variation in redox state of plasma thiol/disulfide couples occurs in various physiologic conditions, including diabetes, chemotherapy, and aging. The present study was designed to determine whether a systematic variation in extracellular thiol/disulfide redox state (E(h)) over a range (0 mV to -150 mV) that occurs in human plasma altered proliferation of cultured cells. Experiments were performed with a human colon carcinoma cell line (Caco2), which grows slowly in the absence of serum and responds to peptide growth factors with increased rate of cell division. The extracellular redox states were established by varying concentrations of cysteine and cystine, maintaining constant pool size in terms of cysteine equivalents. Incorporation of 5-bromo-2-deoxyuridine (BrdU) was used to measure DNA synthesis and was lowest at the most oxidized extracellular E(h) (0 mV). Incorporation increased as a function of redox state, attaining a 100% higher value at the most reduced condition (-150 mV). Addition of insulin-like growth factor-1 (IGF-1) or epidermal growth factor (EGF) increased the rate of BrdU

incorporation at more oxidizing redox conditions (0 to -80 mV) but had no effect at -150 mV. Cellular GSH was not significantly affected by variation in extracellular E(h). In the absence of growth factors, extracellular E(h) values were largely maintained for 24 h. However, IGF-1 or EGF stimulated a change in extracellular redox to values similar to that for cysteine/cystine redox in plasma of young, healthy individuals. The results show that extracellular thiol/disulfide redox state modulates cell proliferation rate and that this control interacts with growth factor signaling apparently independently of cellular glutathione

L38 ANSWER 21 OF 67 MEDLINE ON STN ACCESSION NUMBER: 2002672874 MEDLINE DOCUMENT NUMBER: PubMed ID: 12433666

TITLE: Glutathione and thioredoxin

redox during differentiation in human colon

epithelial (Caco-2) cells.

AUTHOR: Nkabyo Yvonne S; Ziegler Thomas R; Gu Li H; Watson Walter

H; Jones Dean P

CORPORATE SOURCE: Department of Biochemistry, the Graduate Program in

Molecular and Systems Pharmacology, Emory University,

Atlanta, Georgia 30322, USA.

CONTRACT NUMBER: DK-55850 (NIDDK)

ES-009047 (NIEHS) ES-011195 (NIEHS) RR-00039 (NCRR)

SOURCE: American journal of physiology. Gastrointestinal and liver

physiology, (2002 Dec) 283 (6) G1352-9. Journal code: 100901227. ISSN: 0193-1857.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200212

ENTRY DATE: Entered STN: 20021116

Last Updated on STN: 20021218 Entered Medline: 20021213

AB Cellular redox, maintained by the glutathione (GSH) and thioredoxin (Trx) -dependent systems, has been implicated in the regulation of a variety of biological processes. The redox state of the GSH system becomes oxidized when cells are induced to differentiate by chemical agents. The aim of this study was to determine the redox state of cellular GSH/qlutathione disulfide (GSH/GSSG) and Trx as a consequence of progression from proliferation to contact inhibition and spontaneous differentiation in colon carcinoma (Caco-2) cells. Results showed a significant decrease in GSH concentration, accompanied by a 40-mV oxidation of the cellular GSH/GSSG redox state and a 28-mV oxidation of the extracellular cysteine/cystine redox state in association with confluency and increase in differentiation markers. The redox state of Trx did not change. Thus the two central cellular antioxidant and redox-regulating systems (GSH and Trx) were independently controlled. According to the Nernst equation, a 30-mV oxidation is associated with a 10-fold change in the reduced/oxidized ratio of a redox-sensitive dithiol motif. Therefore, the measured 40-mV oxidation of the cellular GSH/GSSG couple or the 28-mVoxidation of the extracellular cysteine/cystine couple should be sufficient to function in signaling or regulation of differentiation in

Caco-2 cells.

L38 ANSWER 22 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2002384053 EMBASE

TITLE: Glutathione catabolism as a signaling mechanism.

AUTHOR: Paolicchi A.; Dominici S.; Pieri L.; Maellaro E.; Pompella

Α.

CORPORATE SOURCE: A. Pompella, Department of Experimental Pathology,

University of Pisa Medical School, Via Roma 55, 56126 Pisa,

Italy. apompella@biomed.unipi.it

SOURCE: Biochemical Pharmacology, (1 Sep 2002) Vol. 64, No. 5-6,

pp. 1027-1035.

Refs: 38

ISSN: 0006-2952 CODEN: BCPCA6

PUBLISHER IDENT.: S 0006-2952(02)01173-5

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20021121

Last Updated on STN: 20021121

AB Glutathione (GSH) is the main intracellular thiol antioxidant, and as such participates in a number of cellular antitoxic and defensive functions. Nevertheless, non-antioxidant functions of GSH have also been decribed, e.g. in modulation of cell proliferation and immune response. Recent studies from our and other laboratories have provided evidence for a third functional aspect of GSH, i.e. the prooxidant roles played by molecular species originating during its catabolism by the membrane ectoenzyme γ -glutamyl transpeptidase (GGT). The reduction of metal ions effected by GSH catabolites is capable to induce redox cycling processes leading to the production of reactive oxygen species (superoxide, hydrogen peroxide), as well as of other free radicals. Through the action of these reactive compounds, GSH catabolism can ultimately lead to oxidative modifications on a variety of molecular targets, involving oxidation and/or S-thiolation of protein thiol groups in the first place. Modulating effects of this kind have been observed on several important, redox -sensitive components of the signal transduction chains, such as cell surface receptors, protein phosphatase activities and transcription factors. Against this background, the prooxidant reactions induced by GSH

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modulation of cellular signal transduction. . COPYRGT. 2002 Elsevier

catabolism appear to represent a novel, as yet unrecognized mechanism for

on STN

AUTHOR ·

ACCESSION NUMBER: 2002172879 EMBASE

TITLE: Dendritic cells generate thiols for T-cell

proliferation.
Keenihan S.H.

CORPORATE SOURCE: keenihan@namru2.med.navy.mil

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SOURCE: Trends in Immunology, (1 May 2002) Vol. 23, No. 5, pp. 233.

Refs: 1

ISSN: 1471-4906 CODEN: TIRMAE

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Note

FILE SEGMENT: 026 Immunology, Serology and Transplantation

037 Drug Literature Index

LANGUAGE: English

ENTRY DATE: Entered STN: 20020530

Last Updated on STN: 20020530 DATA NOT AVAILABLE FOR THIS ACCESSION NUMBER

L38 ANSWER 24 OF 67 MEDLINE on STN DUPLICATE 11

ACCESSION NUMBER: 2002141305 MEDLINE DOCUMENT NUMBER: PubMed ID: 11849044

TITLE: Inhibition of cell proliferation and AP-1

activity by acrolein in human A549 lung

adenocarcinoma cells due to thiol
imbalance and covalent modifications.

AUTHOR: Biswal Shyam; Acquaah-Mensah George; Datta Kaushik; Wu

Xuli; Kehrer James P

CORPORATE SOURCE: Bloomberg School of Public Health, Johns Hopkins

University, Department of Environmental Health Sciences, Division of Toxicological Sciences, Baltimore, Maryland

21205-2179, USA.. sbiswal@jhsph.edu

CONTRACT NUMBER: ES09791 (NIEHS)

F32 ES05896 (NIEHS)

SOURCE: Chemical research in toxicology, (2002 Feb) 15 (2) 180-6.

Journal code: 8807448. ISSN: 0893-228X.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200204

ENTRY DATE: Entered STN: 20020307

Last Updated on STN: 20020424 Entered Medline: 20020423

AB Acrolein, a reactive alpha, beta-unsaturated aldehyde, is a common environmental pollutant, a metabolite of the anticancer drug cyclophosphamide, and a byproduct of lipid peroxidation. An increase in acrolein production has been proposed as a marker for Alzheimer's disease, diabetic glomerular lesions, and atherosclerosis. Acrolein is a potent inhibitor of cell proliferation at nonlethal doses and may act through effects on redox-regulated transcription factors. We previously reported that NF-kappaB activation is inhibited by acrolein in the A549 lung adenocarcinoma cell line in an IkappaB-independent manner [Horton et al. (1999) J. Biol. Chemical 274, 9200-9206]. current data demonstrate that AP-1 activation in A549 cells is decreased by 26 and 50% at 0.5 and 1 h, respectively, after exposure to 50 fmol/cell (a nonlethal dose) of acrolein. Inhibition of AP-1 activation also occurred following treatment with buthionine sulfoximine to deplete glutathione to the same extent as seen with acrolein. c-jun antisense treatments depressed c-jun protein below detectable levels at 4 h and inhibited cell proliferation (as assessed by [(3)H]thymidine incorporation) by 80%. Immunoprecipitation of c-jun protein after treating A549 cells with acrolein revealed the presence of a lysine-acrolein adduct. There was, however, no effect of acrolein on c-jun N-terminal kinase activity or c-jun phosphorylation. These data indicate that the inhibition of cell proliferation induced by acrolein correlates with the depletion of glutathione as well as the inhibition of AP-1 activation. AP-1 activation is likely affected both through changes in cellular thiol redox balance and by covalent modification of acrolein to c-jun, but not through effects on c-jun phosphorylation.

L38 ANSWER 25 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 2002421555 EMBASE

TITLE: Glutathione and thioredoxin

redox during differentiation in human colon

epithelial (Caco-2) cells.

AUTHOR: Nkabyo Y.S.; Ziegler T.R.; Gu L.H.; Watson W.H.; Jones D.P.

CORPORATE SOURCE: D.P. Jones, Dept. of Biochemistry, Rollins Research Center,

Emory University, 1510 Clifton Rd. NE, Atlanta, GA 30322,

United States. dpjones@emory.edu

SOURCE: American Journal of Physiology - Gastrointestinal and Liver

Physiology, (1 Dec 2002) Vol. 283, No. 6 46-6, pp.

G1352-G1359. Refs: 39

ISSN: 0193-1857 CODEN: APGPDF

COUNTRY: United States
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 002 Physiology

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 20021212

Last Updated on STN: 20021212

AB Cellular redox, maintained by the glutathione (GSH) and thioredoxin (Trx)-dependent systems, has been implicated in
the regulation of a variety of biological processes. The redox
state of the GSH system becomes oxidized when cells are induced to
differentiate by chemical agents. The aim of this study was to determine
the redox state of cellular GSH/glutathione disulfide
(GSH/GSSG) and Trx as a consequence of progression from
proliferation to contact inhibition and spontaneous
differentiation in colon carcinoma (Caco-2) cells. Results
showed a significant decrease in GSH concentration, accompanied by a 40-mV

showed a significant decrease in GSH concentration, accompanied by a 40-mV oxidation of the cellular GSH/GSSG redox state and a 28-mV oxidation of the extracellular cysteine/cystine redox state in association with confluency and increase in differentiation markers. The redox state of Trx did not change. Thus the two central cellular antioxidant and redox-regulating systems (GSH and Trx) were independently controlled. According to the Nernst equation, a 30-mV oxidation is associated with a 10-fold change in the reduced/oxidized ratio of a redox-sensitive dithiol motif. Therefore, the measured 40-mV oxidation of the cellular GSH/GSSG couple or the 28-mV oxidation of the extracellular cysteine/cystine couple should be

oxidation of the extracellular cysteine/cystine couple should be sufficient to function in signaling or regulation of differentiation in Caco-2 cells.

L38 ANSWER 26 OF 67 MEDLINE on STN DUPLICATE 12

ACCESSION NUMBER: 2001287563 MEDLINE DOCUMENT NUMBER: PubMed ID: 11278531

TITLE: Cyclopentenone prostaglandins as potential inducers of

intracellular oxidative stress.

AUTHOR: Kondo M; Oya-Ito T; Kumagai T; Osawa T; Uchida K

CORPORATE SOURCE: Laboratory of Food and Biodynamics, Graduate School of

Bioagricultural Sciences, Nagoya University, Nagoya

464-8601, Japan.

SOURCE: Journal of biological chemistry, (2001 Apr 13) 276 (15)

12076-83. Electronic Publication: 2001-01-12.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200105

ENTRY DATE: Entered STN: 20010529

Last Updated on STN: 20030105 Entered Medline: 20010524

AB In the present study, we find that cyclopentenone prostaglandins (PGs) of the J(2) series, naturally occurring derivatives of PGD(2), are potential inducers of intracellular oxidative stress that mediates cell degeneration. Based on an extensive screening of diverse chemical agents on induction of intracellular production of reactive oxygen species (ROS), we found that the cyclopentenone PGs, such as PGA(2), PGJ(2), Delta(12)-PGJ(2), and 15-deoxy-Delta(12,14)-PGJ(2), showed the most potent pro-oxidant effect on SH-SY5Y human neuroblastoma cells. As the intracellular events that mediate the PG cytotoxicity, we observed (i) the cellular redox alteration represented by depletion of antioxidant defenses, such as glutathione and glutathione peroxidase; (ii) a transient decrease in the mitochondrial membrane potential (Deltapsi); (iii) the production of protein-bound lipid peroxidation products, such as acrolein and 4-hydroxy-2-nonenal; and (iv) the accumulation of ubiquitinated proteins. These events correlated well with the reduction in cell viability. In addition, the thiol compound, N-acetylcysteine , could significantly inhibit the PG-induced ROS production, thereby preventing cytotoxicity, suggesting that the redox alteration is closely related to the pro-oxidant effect of cyclopentenone PGs. strikingly, the lipid peroxidation end products, acrolein and 4-hydroxy-2-nonenal, detected in the PG-treated cells potently induced the ROS production, which was accompanied by the accumulation of ubiquitinated proteins and cell death, suggesting that the membrane lipid peroxidation products may represent one of the causative factors that potentiate the cytotoxic effect of cyclopentenone PGs by accelerating intracellular oxidative stress. These data suggest that the intracellular oxidative stress, represented by ROS production/lipid peroxidation and redox alteration, may underlie the well documented biological effects, such as antiproliferative and antitumor activities, of cyclopentenone PGs.

L38 ANSWER 27 OF 67 MEDLINE on STN DUPLICATE 13

ACCESSION NUMBER: 2001111315 MEDLINE DOCUMENT NUMBER: PubMed ID: 11123355

TITLE: Cellular thiols and reactive oxygen species in

drug-induced apoptosis.

AUTHOR: Davis W Jr; Ronai Z; Tew K D

CORPORATE SOURCE: Department of Pharmacology, Fox Chase Cancer Center,

Philadelphia, Pennsylvania, USA.

CONTRACT NUMBER: CA06927 (NCI)

CA85660 (NCI) RR05539 (NCRR)

SOURCE: Journal of pharmacology and experimental therapeutics,

(2001 Jan) 296 (1) 1-6. Ref: 43

Journal code: 0376362. ISSN: 0022-3565.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200102

ENTRY DATE: Entered STN: 20010322

Last Updated on STN: 20010322 Entered Medline: 20010202

In higher eukaryotes, reactive oxygen species (ROS) are generated during respiration in mitochondria in the course of reduction of molecular oxygen as well as by distinct enzyme systems. ROS have been implicated in the regulation of diverse cellular functions including defense against pathogens, intracellular signaling, transcriptional activation, proliferation, and apoptosis. The reduction-oxidation (redox) state of the cell is primarily a consequence of the precise balance between the levels of ROS and endogenous thiol buffers present in the cell, such as glutathione and thioredoxin, which protect cells from oxidative damage. Dramatic elevation of ROS, exceeding compensatory changes in the level of the endogenous thiol buffers, may result in the sustained activation of signaling

, which protect cells from oxidative damage. Dramatic elevation of ROS, exceeding compensatory changes in the level of the endogenous thiol buffers, may result in the sustained activation of signaling pathways and expression of genes that induce apoptosis in affected cells. Many cytotoxic drugs function selectively to kill cancer cells by the abrogation of proliferative signals, leading to cell death, and numerous reports have demonstrated that ROS are generated following treatment with these drugs. In this review, we will summarize recent contributions to our understanding of the importance of cytotoxic drug-induced modulation of cellular redox status for signaling and transcription leading to activation of apoptotic effector mechanisms.

L38 ANSWER 28 OF 67 MEDLINE on STN DUPLICATE 14

ACCESSION NUMBER: 2001040936 MEDLINE DOCUMENT NUMBER: PubMed ID: 11029606

TITLE: Virological and immunological effects of antioxidant

treatment in patients with HIV infection.

AUTHOR: Muller F; Svardal A M; Nordoy I; Berge R K; Aukrust P;

Froland S S

CORPORATE SOURCE: University of Oslo, The National Hospital, Rikshospitalet,

Oslo, Norway.. fredrik.muller@labmed.uio.no

SOURCE: European journal of clinical investigation, (2000 Oct) 30

(10) 905-14.

Journal code: 0245331. ISSN: 0014-2972.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: (CLINICAL TRIAL)

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200012

ENTRY DATE: Entered STN: 20010322

Last Updated on STN: 20010322 Entered Medline: 20001207

AB BACKGROUND: Intracellular oxidative stress in CD4+ lymphocytes due to disturbed glutathione homeostasis may lead to impaired lymphocyte functions and enhanced HIV replication in patients with HIV infection, especially in those with advanced immunodeficiency. The aim of the present study was to assess whether short-term, high-dose antioxidant treatment might have effects on immunological and virological parameters in patients with HIV infection. MATERIALS AND METHODS: In this pilot study, we examined virological and immunological effects of antioxidant combination treatment for 6 days with high doses of N-acetylcysteine (NAC) and vitamin C in 8 patients with HIV infection. The following were assayed before, during and after antioxidant treatment: HIV RNA plasma levels; numbers of CD4+, CD8+, and CD14+ leukocytes in blood; plasma thiols; intracellular

glutathione redox status in CD4+ lymphocytes and CD14+ monocytes; lymphocyte proliferation; lymphocyte apoptosis and plasma levels of tumour necrosis factor (TNF)alpha; soluble TNF receptors and neopterin in plasma. RESULTS: No significant changes in HIV RNA plasma levels or CD4+ lymphocyte counts in blood were noted during antioxidant treatment in the patient group. However, in the 5 patients with the most advanced immunodeficiency (CD4+ lymphocyte counts < 200 x 106 L(-1)), a significant rise in CD4+ lymphocyte count, a reduction in HIV RNA plasma level of 0.8 log, an enhanced lymphocyte proliferation and an increased level of intracellular glutathione in CD4+ lymphocytes were found. No change in lymphocyte apoptosis was noted. CONCLUSIONS: Short-term, high-dose combination treatment with NAC and vitamin C in patients with HIV infection and advanced immunodeficiency lead to immunological and virological effects that might be of therapeutic value.

L38 ANSWER 29 OF 67 MEDLINE on STN DUPLICATE 15

ACCESSION NUMBER: 1999418846 MEDLINE DOCUMENT NUMBER: PubMed ID: 10490284

TITLE: Redox modulation of cell surface protein

thiols in U937 lymphoma cells: the role of

gamma-glutamyl transpeptidase-dependent H2O2 production and

S-thiolation.

AUTHOR: Dominici S; Valentini M; Maellaro E; Del Bello B; Paolicchi

A; Lorenzini E; Tongiani R; Comporti M; Pompella A

CORPORATE SOURCE: Institute of General Pathology, University of Siena, Italy.

SOURCE: Free radical biology & medicine, (1999 Sep) 27 (5-6)

623-35.

Journal code: 8709159. ISSN: 0891-5849.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199910

ENTRY DATE: Entered STN: 19991101

Last Updated on STN: 19991101 Entered Medline: 19991019

AB The expression of gamma-glutamyl transpeptidase (GGT), a plasma membrane ectoenzyme involved in the metabolism of extracellular reduced glutathione (GSH), is a marker of neoplastic progression in several experimental models, and occurs in a number of human malignant neoplasms and their metastases. Because it favors the supply of precursors for the synthesis of GSH, GGT expression has been interpreted as a member in cellular antioxidant defense systems. However, thiol metabolites generated at the cell surface during GGT activity can induce prooxidant reactions, leading to production of free radical oxidant species. The present study was designed to characterize the prooxidant reactions occurring during GGT ectoactivity, and their possible effects on the thiol redox status of proteins of the cell surface. Results indicate that: (i) in U937 cells, expressing significant amounts of membrane-bound GGT, GGT-mediated metabolism of GSH is coupled with the extracellular production of hydrogen peroxide; (ii) GGT activity also results in decreased levels of protein thiols at the cell surface; (iii) GGT-dependent decrease in protein thiols is due to sulfhydryl oxidation and protein Sthiolation reactions; and (iv) GGT irreversible inhibition by acivicin is sufficient to produce an increase of protein thiols at the cell surface. Membrane receptors and transcription factors have been shown to possess critical thiols involved in

the transduction of **proliferative** signals. Furthermore, it was suggested that S-**thiolation** of cellular proteins may represent a mechanism for protection of vulnerable **thiols** against irreversible damage by prooxidant agents. Thus, the findings reported here provide additional explanations for the envisaged role played by membrane-bound GGT activity in the **proliferative** attitude of malignant cells and their resistance to prooxidant drugs and radiation therapy.

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on STN

ACCESSION NUMBER: 1999010471 EMBASE

TITLE: Changes in glutathione status and the antioxidant

system in blood and in cancer cells associate

with tumour growth in vivo.

AUTHOR: Navarro J.; Obrador E.; Carretero J.; Petschen I.; Avino

J.; Perez P.; Estrela J.M.

CORPORATE SOURCE: Dr. J.M. Estrela, Departamento de Fisiologia, Facultad de

Medicina, Av. Biasco Ibanez 17, 46010 Valen, Spain.

jose.mlestrela@uv.es

SOURCE: Free Radical Biology and Medicine, (1999) Vol.

26, No. 3-4, pp. 410-415.

Refs: 47

ISSN: 0891-5849 CODEN: FRBMEH

PUBLISHER IDENT.: S 0891-5849(98)00213-5

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

016 Cancer

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19990204

Last Updated on STN: 19990204

AB The relationship among cancer growth, the glutathione redox cycle and the antioxidant system was studied in blood and in

tumour cells. During cancer growth, the

glutathione redox status (GSH/GSSG) decreases in blood

of Ehrlich ascites tumour-bearing mice. This effect is mainly

due to an increase in GSSG levels. Two reasons may explain the increase in blood GSSG: (a) the increase in peroxide production by the

tumour that, in addition to changes affecting the

glutathione-related and the antioxidant enzyme activities, can

lead to GSH oxidation within the red blood cells; and (b) an increase of GSSG release from different tissues into the blood. GSH and peroxide

levels are higher in the tumour cells when they

proliferate actively, however GSSG levels remain constant during turnout growth in mice. These changes associate with low levels of lipid

peroxidation in plasma, blood and the tumour cells. The

GSH/GSSG ratio in blood also decreases in patients bearing breast or colon

cancers and, as it occurs in tumour-bearing mice, this

change associates with higher GSSG levels, especially in advanced stages

of cancer progression. Our results indicate that determination of clutathions status and oxidative stress-related parameters in

of glutathione status and oxidative stress-related parameters in

blood may help to orientate cancer therapy in humans.

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on STN

ACCESSION NUMBER: 1999213264 EMBASE

TITLE: Induction of thioredoxin by oxidative stress and

overexpression of thioredoxin in lung

cancer tissue.

AUTHOR: Jang Hoon Lee; Hyung Jung Kim; Chul Min Ahn; Sung Kyu Kim;

Won Young Lee

CORPORATE SOURCE: Dr. J.H. Lee, Department of Internal Medicine, Yonsei Univ.

College of Medicine, Seoul, Korea, Republic of Tuberculosis and Pespiratory Diseases (1999)

Tuberculosis and Respiratory Diseases, (1999)

Vol. 46, No. 3, pp. 327-337.

Refs: 32

ISSN: 0378-0066 CODEN: KHCHAM

COUNTRY: Korea, Republic of DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

Ol5 Chest Diseases, Thoracic Surgery and Tuberculosis

037 Drug Literature Index

LANGUAGE: Korean

SOURCE:

SUMMARY LANGUAGE: English; Korean ENTRY DATE: Entered STN: 19990708

Last Updated on STN: 19990708

Background: Reactive oxygen species are involved in multi-stage process of AB carcinogenesis. The most of cancer cell lines and cancer cells in tumor tissue produce reactive oxygen species and on the other hand, the activities of catalase, Mn- and CuZn-superoxide dismutase in tumor cells are usually low. persistent oxidative stress in tumor tissue facilitates tumor invasion and metastasis. 12-kDa thioredoxin, which regulates the intracellular redox potential with glutathione and glutaredoxin is involved in cell activation, proliferation, differentiation and redox -mediated apoptosis. It is also purified as 14-kDa and 10-kDa eosinophilic cytotoxic enhancing factor(ECEF) from human histiocytic cell(U937) and 10-kDa ECEF has more than 20 times eosinophilic stimulation activity than 14-kDa ECEF. It has been reported that adult T-cell leukemia, squamous cell carcinoma of uterine cervix, and hepatocellular carcinoma show increased amounts of human thioredoxin and thioredoxin mRNA is increased in lung cancer. In this study, we investigated the expression of conventional antioxidant enzymes such as catalase, CuZn-SOD, and glutathione peroxidase and thioredoxin in lung cancer tissue compared to adjacent normal lung tissue and the induction of thioredoxin in macrophage cells after treatment of oxidative stress and endotoxin. Methods: We measured the amount of conventional antioxidant enzymes such as catalase, CuZn-SOD, and glutathione peroxidase and thioredoxin in lung cancer tissue compared to adjacent normal lung tissue by immunoblot analysts and the induction of thioredoxin in mouse monocyte- macrophage cells(RAW 264.7) by treatment of 5 μ M menadione and 1 µg/ml endotoxin. Results: On immunoblot analysis, the expression of 12-kDa thioredoxin was increased in lung cancer tissue compared to paired normal lung tissue, but the expression of catalase and CuZn-SOD were decreased in lung cancer tissue compared to paired normal tissue and the expression of glutathione peroxidase in lung cancer was variable. The expression of truncated thioredoxin was also increased in lung cancer . When mouse monocyte- macrophage cells were treated with 5 μM menadione and 1 $\mu g/ml$ endotoxin, the expression of thioredoxin

was peaked at 12 hrs and sustained to 48 hrs. Conclusion: In contrast with other conventional antioxidants, the expression of 12-kDa and

truncated thioredoxin in lung cancer were increased and it is closely associated with persistent oxidative stress in tumor microenvironment. Considering especially the biological functions of truncated thioredoxin, the increased amount of truncated thioredoxin has significant role in tumor growth through cell proliferation.

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on STN

ACCESSION NUMBER: 1999419649 EMBASE

Increased glutathione synthesis associated with TITLE:

platelet-derived growth factor stimulation of NIH3T3

fibroblasts.

Iantomasi T.; Favilli F.; Degl'Innocenti D.; Vincenzini AUTHOR:

M.T.

CORPORATE SOURCE: M.T. Vincenzini, Department of Biochemical Sciences,

University of Florence, viale Morgagni 50, 50134 Florence,

Italy. vincenzini@cesit1.unifi.it

SOURCE: Biochimica et Biophysica Acta - Molecular Cell Research, (

1999) Vol. 1452, No. 3, pp. 303-312.

Refs: 46

ISSN: 0167-4889 CODEN: BAMRDP

PUBLISHER IDENT.: S 0167-4889(99)00142-1

COUNTRY:

Netherlands Journal; Article

DOCUMENT TYPE:

FILE SEGMENT: LANGUAGE:

029 Clinical Biochemistry

SUMMARY LANGUAGE:

English English

ENTRY DATE:

Entered STN: 19991229

Last Updated on STN: 19991229

AB Previous data show a relation between GSH content and proliferation of normal and tumour cells. We recently demonstrated a specific involvement of GSH in the autophosphorylation activity of the platelet-derived growth factor (PDGF) receptor in NIH3T3 fibroblasts. In this study we demonstrate that the stimulation by PDGF of serum-starved NIH3T3 cells increases cellular GSH content, while no change in oxidized GSH content was measured. Experiments performed with actinomycin, cycloheximide and buthionine sulfoximide, a specific inhibitor of the rate-limiting enzyme of the de novo synthesis of GSH γ -glutamylcysteine synthetase (γ -GCS), confirm PDGF induction of GSH synthesis. These results provide the first demonstration that PDGF mediated transduction signals seem strictly related to mechanisms involved in the increase of γ -GCS activity associated with increased γ -GCS heavy subunit mRNA levels. In fact, serum and epidermal growth factor (EGF) stimulation of quiescent NIH3T3 and NIH3T3, which overexpress EGF receptor, does not affect GSH content or its synthesis. These data may be related to a possible GSH role in the redox regulation of cell proliferation mediated by PDGF.

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on STN

CORPORATE SOURCE:

ACCESSION NUMBER: 1999396799 EMBASE

Redox regulation of TNF signaling. TITLE

AUTHOR: Goossens V.; De Vos K.; Vercammen D.; Steemans M.;

> Vancompernolle K.; Fiers W.; Vandenabeele P.; Grooten J. J. Grooten, Department of Molecular Biology, Flanders

Interuniversity, Institute for Biotechnology, K.L.

Ledeganckstraat 35, B-9000 Gent, Belgium

SOURCE: BioFactors, (1999) Vol. 10, No. 2-3, pp. 145-156. Refs: 37

ISSN: 0951-6433 CODEN: BIFAEU

COUNTRY: Netherlands

DOCUMENT TYPE: Journal; Conference Article FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19991202

Last Updated on STN: 19991202

TNF is produced during inflammation and induces, among other activities, AB cell death in sensitive tumour cells. We previously reported an increased generation of ROS in TNF-treated L929 fibrosarcoma cells prior to cell death. These ROS are of mitochondrial origin and participate in the cell death process. Presently, we focus on the identification of parameters that control ROS production and subsequent cytotoxicity. From the cytotoxic properties and susceptibility to scavenging of TNF-induced ROS as compared to pro-oxidant- induced ROS we conclude that TNF-mediated ROS generation and their lethal action are confined to the inner mitochondrial membrane. Oxidative substrates, electron-transport inhibitors, glutathione and thiol-reactive agents but also caspase inhibitors modulate TNF-induced ROS production and imply the existence of a negative regulator of ROS production. Inactivation of this regulator by a TNF-induced reduction of NAD(P)H levels and/or formation of intraprotein disulfides would be responsible for ROS generation.

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on STN

ACCESSION NUMBER: 1998215830 EMBASE

TITLE: Disruption of redox homeostasis in the

transforming growth factor- α/c - myc transgenic mouse

model of accelerated hepatocarcinogenesis.

AUTHOR: Factor V.M.; Kiss A.; Woitach J.T.; Wirth P.J.;

Thorgeirsson S.S.

CORPORATE SOURCE: S.S. Thorgeirsson, National Cancer Institute, NIH, Bldg.

37, 37 Convent Drive MSC4255, Bethesda, MD 20892-4255,

United States

SOURCE: Journal of Biological Chemistry, (19 Jun 1998)

Vol. 273, No. 25, pp. 15846-15853.

Refs: 81

ISSN: 0021-9258 CODEN: JBCHA3

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19980806

Last Updated on STN: 19980806

AB In previous studies we have demonstrated that transforming growth factor (TGF)-α/c-myc double transgenic mice exhibit an enhanced rate of cell proliferation, accumulate extensive DNA damage, and develop multiple liver tumors between 4 and 8 months of age. To clarify the biochemical events that may be responsible for the genotoxic and carcinogenic effects observed in this transgenic model, several parameters of redox homeostasis in the liver were examined prior to development of hepatic tumors. By 2 months of age, production of reactive oxygen species, determined by the peroxidation-sensitive fluorescent dye, 2',7'-dichlorofluorescin diacetate, was significantly elevated in TGF-α/c-myc transgenic hepatocytes versus either wild type or c-myc single transgenic cells, and occurred in

parallel with an increase in lipid peroxidation. Concomitantly with a rise in oxidant levels, antioxidant defenses were decreased, including total glutathione content and the activity of glutathione peroxidase, whereas thioredoxin reductase activity was not changed. However, hepatic tumors which developed in TGF- α/c -myc mice exhibited an increase in thioredoxin reductase activity and a very low activity of glutathione peroxidase. Furthermore, specific deletions were detected in mtDNA as early as 5 weeks of age in the transgenic mice. These data provide experimental evidence that co-expression of TGF- α and c- myc transgenes in mouse liver promotes overproduction of reactive oxygen species and thus creates an oxidative stress environment. This phenomenon may account for the massive DNA damage and acceleration of hepatocarcinogenesis observed in the TGF- α/c -myc mouse model.

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on STN

ACCESSION NUMBER: 1998215763 EMBASE

TITLE: Reversible inactivation of protein-tyrosine phosphatase 1B

in A431 cells stimulated with epidermal growth factor.

AUTHOR: Lee S.-R.; Kwon K.-S.; Kim S.-R.; Rhee S.G.

CORPORATE SOURCE: S.G. Rhee, Bldg. 3, National Institutes of Health,

Bethesda, MD 20892, United States. sgrhee@helix.nih.gov

SOURCE: Journal of Biological Chemistry, (19 Jun 1998)

Vol. 273, No. 25, pp. 15366-15372.

Refs: 54

ISSN: 0021-9258 CODEN: JBCHA3

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19980806

Last Updated on STN: 19980806

Stimulation of various cells with growth factors results in a transient AB increase in the intracellular concentration of H2O2 that is required for growth factor-induced protein tyrosine phosphorylation. The effect of H202 produced in response to epidermal growth factor (EGF) on the activity of protein-tyrosine phosphatase 1B (PTP1B) was investigated in A431 human epidermoid carcinoma cells. H2O2 inactivated recombinant PTP1B in vitro by oxidizing its catalytic site cysteine, most likely to sulfenic acid. The oxidized enzyme was reactivated more effectively by thioredoxin than by glutaredoxin or glutathione at their physiological concentrations. Oxidation by H2O2 prevented modification of the catalytic cysteine of PTP1B by iodoacetic acid, suggesting that it should be possible to monitor the oxidation state of PTP1B in cells by measuring the incorporation of radioactivity into the enzyme after lysis of the cells in the presence of radiolabeled iodoacetic acid. The amount of such radioactivity associated with PTP1B immunoprecipitated from A431 cells that had been stimulated with EGF for 10 min was 27% less than that associated with PTP1B from unstimulated cells. The amount of iodoacetic acid-derived radioactivity associated with PTP1B reached a minimum 10 min after stimulation of cells with EGF and returned to base line values by 40 min, suggesting that the oxidation of PTP1B is reversible in cells. These results indicate that the activation of a receptor tyrosine kinase by binding of the corresponding growth factor may not be sufficient to increase the steady state level of protein tyrosine phosphorylation in cells and that concurrent inhibition

of protein-tyrosine phosphatases by H2O2 might also be required.

L38 ANSWER 36 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 1998182919 EMBASE

TITLE: Antioxidants reduce cyclooxygenase-2 expression,

prostaglandin production, and proliferation in

colorectal cancer cells.

AUTHOR: Chinery R.; Beauchamp R.D.; Shyr Y.; Kirkland S.C.; Coffey

R.J.; Morrow J.D.

CORPORATE SOURCE: R.J. Coffey, G1 Cancer Program, CC-2218 Medical Center

North, Vanderbilt University Medical Center, Nashville, TN 37232-2583, United States. coffeyrj@ctrvax.vanderbilt.edu

SOURCE: Cancer Research, (1 Jun 1998) Vol. 58, No. 11,

pp. 2323-2327.

Refs: 23

ISSN: 0008-5472 CODEN: CNREA8

COUNTRY: United States
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 016 Cancer

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19980727

Last Updated on STN: 19980727

AB Increased expression of cyclooxygenase (COX) and overproduction of prostaglandins (PGs) have been implicated in the development and progression of colorectal cancer (CRC). Recent observations suggest that reactive oxygen intermediates play a role in tumor cell growth regulation and expression of the inducible COX, COX-2. We therefore evaluated the effects of various antioxidants on COX expression and cellular growth in the human CRC cell line HCA-7. The antioxidants pyrrolidinedithiocarbamate (PDTC), Nacetylcysteine, 6-hydroxy-2,5,7,8-tetramethylchroman-2-carboxylic acid (Trolox), and U74006 decreased PG production, intracellular redox status, and cellular growth in a concentration-dependent manner. The decrease in cellular growth was associated with the induction of apoptosis. Unlike the selective COX inhibitors 1-[(4methylsulfonyl)phenyl]-3-trifluoromethyl-5-[(4-fluoro)phenyl]pyrazole (SC 58125) and (2-cyclohexyloxy-4- nitrophenyl)methanesulfonamide (NS 398) that inhibit COX-2 catalytic activity, these antioxidants decreased COX-2 expression at the transcriptional level. Combined treatment of HCA-7 cells with PDTC and SC 58125 resulted in an additive decrease in PG levels and anchorage-dependent and -independent growth. Furthermore, whereas antioxidants or SC 58125 reduced tumor growth in vivo, coadministration of PDTC and SC 58125 resulted in actual tumor regression. These results suggest that combined therapy with NSAIDs and antioxidants might be useful in the prevention and/or treatment of CRC.

L38 ANSWER 37 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC: ALL RIGHTS RESERVED. on STN

ACCESSION NUMBER: 1998121794 EMBASE

CORPORATE SOURCE:

TITLE: Mechanisms of inhibition of the thioredoxin

growth factor system by antitumor 2-imidazolyl

disulfides.

AUTHOR: Kirkpatrick D.L.; Kuperus M.; Dowdeswell M.; Potier N.;

Donald L.J.; Kunkel M.; Berggren M.; Angulo M.; Powis G. D.L. Kirkpatrick, Department of Chemistry, University of

Regina, Regina, Sask. S4S 0A2, Canada.

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SOURCE: Biochemical Pharmacology, (1 Apr 1998) Vol. 55,

No. 7, pp. 987-994.

Refs: 26

ISSN: 0006-2952 CODEN: BCPCA6

PUBLISHER IDENT.: S 0006-2952(97)00597-2

COUNTRY: United States
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 016 Cancer

029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19980507

Last Updated on STN: 19980507

AB The interactions of a series of 2-imidazolyl disulfide antitumor compounds with the thioredoxin reductase (TR)/

thioredoxin (hTrx) redox system have been studied. Disulfides III-2 (n-butyl 2-mercaptoimidazolyl disulfide) and VI-2 (ethyl 2-mercaptoimidazolyl disulfide) were substrates for reduction by TR with K(m) values of 43 and 48 μM. Disulfides IV-2 (1-methylpropyl 2-mercaptoimidazolyl disulfide) and DLK-36 (benzyl 2-mercaptoimidazolyl disulfide) were competitive inhibitors of the reduction of hTrx by TR with K(i) values of 31 μM . None of the disulfides were substrates for reduction by human glutathione reductase. The disulfides caused reversible thioalkylation of hTrx at the redox catalytic site as shown by the fact that there was no thioalkylation of a mutant hTrx where both the catalytic site Cys32 and Cys35 residues were replaced by Ser. In addition, the disulfides caused a slower irreversible inactivation of hTrx as a substrate for reduction by TR, with half-lives for III-2 of 30 min, for IV-2 of 4 hr, and for IX-2 (t-butyl 2-mercaptoimidazolyl disulfide) of 24 hr. This irreversible inactivation of hTrx occurred at concentrations of the disulfides an order of magnitude below those that inhibited TR, and involved the Cys73 of hTrx, which is outside the conserved redox catalytic site, as shown by the resistance to inactivation of a mutant hTrx where Cys73 was replaced by Ser. Electrophoretic and mass spectral analyses of the products of the reaction between the disulfides and hTrx show that modification of 1-3 Cys residues of the protein occurred in a concentration-dependent fashion. The disulfides inhibited the hTrx dependent proliferation of MCF-7 breast cancer cells with IC50 values for III-2 and IV-2 of 0.2 and 1.2 μ M, respectively. The results show that although the catalytic sites of TR and hTrx are reversibly inhibited by the 2-imidazolyl disulfides, it is the irreversible thioalkylation of Cys73 of hTrx by the disulfides chat most probably accounts for the inhibition of thioredoxin dependent

L38 ANSWER 38 OF 67 MEDLINE on STN DUPLICATE 16

ACCESSION NUMBER: 1998138892 MEDLINE DOCUMENT NUMBER: PubMed ID: 9518260

cell growth by the disulfides.

TITLE: Thiol redox modulation of tumor

necrosis factor-alpha responsiveness in cultured

AIDS-related Kaposi's sarcoma cells.

AUTHOR: Mallery S R; Landwehr D J; Ness G M; Clark Y M; Hohl C M CORPORATE SOURCE: Departments of Oral Surgery and Pathology, Colleges of

Dentistry and Medicine, Ohio State University, Columbus

43210, USA.

CONTRACT NUMBER: CA UO1 66531 (NCI)

DE RO1 12183 (NIDCR)

R01 48547

SOURCE: Journal of cellular biochemistry, (1998 Mar 1) 68 (3)

339-54

Journal code: 8205768. ISSN: 0730-2312.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals; AIDS

ENTRY MONTH: 199803

ENTRY DATE: Entered STN: 19980407

Last Updated on STN: 20000303 Entered Medline: 19980325

AΒ Both clinical and experimental evidence indicates that AIDS-related Kaposi's sarcoma (AIDS-KS) has a multifactorial pathogenesis with factors such as HIV viral load, latent virus induction, and opportunistic infections contributing to disease progression. However, a consistent feature that unites these apparently diverse putative etiologic agents is sustained serum elevations of pro-inflammatory cytokines such as tumor necrosis factor-alpha (TNF-alpha). While virtually every cell responds to TNF-alpha with gene activation, the extent of TNF-alpha-mediated cellular signaling is regulated by a delicate balance between signal activation and signal arresting events. Reactive oxygen intermediates (ROI), which are generated as a consequence of TNF-alpha membrane interaction, are part of this TNF-alpha-initiated cellular activation cascade. Previous studies in our laboratory have shown that AIDS-KS cells possess impaired oxygen intermediate scavenging capacities, thereby establishing conditions permissive for the intracellular retention of ROI. In this study, we used cellular capacity to upregulate the cytoprotective enzyme superoxide dismutase (SOD) to address the extent of cellular response to TNF-alpha. Concurrent with the SOD analyses, nucleotide profiles were obtained to assess cellular bioenergetic responses during TNF-alpha challenge. Proliferative growth levels of mitochondrial (Mn)SOD activities showed an activity spectrum ranging from lowest activity in AIDS-KS cells, to intermediate levels in matched, nonlesional cells from the AIDS-KS donors, to highest activities in HIV normal fibroblasts. In contrast, following TNF-alpha challenge, the AIDS-KS and KS donor nonlesional cells showed a 11.89- and 5.86-fold respective increase in MnSOD activity, while the normal fibroblasts demonstrated a 1.35-fold decrease. Subsequent thiol redox modulation studies showed that only the normal fibroblast cultures showed a potentiation of TNF-alpha-mediated MnSOD upregulation following GSH depletion. In addition, provision of the GSH precursor, N-acetylcysteine during TNF-alpha challenge only diminished MnSOD activity and mitochondrial compartmentalization in the AIDS-KS cells, a finding that likely reflects the lower levels of reduced thiols in this cellular population. Our data, which show that a perturbation in their cellular thiol redox status accentuates AIDS-KS cellular responsiveness to TNF-alpha, suggest a biochemical rationale for the recognized TNF-alpha AIDS-KS clinical correlation.

L38 ANSWER 39 OF 67 MEDLINE on STN DUPLICATE 17

ACCESSION NUMBER: 1998049564 MEDLINE DOCUMENT NUMBER: PubMed ID: 9388242

TITLE: Regulatory role for a novel human thioredoxin

peroxidase in NF-kappaB activation.

AUTHOR: Jin D Y; Chae H Z; Rhee S G; Jeang K T

CORPORATE SOURCE: Laboratory of Molecular Microbiology, NIAID, National

Institutes of Health, Bethesda, Maryland 20892, USA.

SOURCE: Journal of biological chemistry, (1997 Dec 5) 272 (49)

30952-61.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals; AIDS

OTHER SOURCE: GENBANK-U25182

ENTRY MONTH: 199801

ENTRY DATE: Entered STN: 19980122

Last Updated on STN: 20030204 Entered Medline: 19980108

AΒ Reduction-oxidation (redox) plays a critical role in NF-kappaB activation. Diverse stimuli appear to utilize reactive oxygen species (e.g. hydrogen peroxide) as common effectors for activating NF-kappaB. Antioxidants govern intracellular redox status, and many such molecules can reduce H2O2. However, functionally, it does appear that different antioxidants are variously selective for redox regulation of certain transcription factors such as NF-kappaB. For NF-kappaB, thioredoxin has been described to be a more potent antioxidant than either glutathione or Nacetylcysteine. Thioredoxin peroxidase is the immediate enzyme that links reduction of H2O2 to thioredoxin. Several putative human thioredoxin peroxidases have been identified using recursive sequence searches/alignments with yeast or prokaryotic enzymes. None has been characterized in detail for intracellular function(s). Here, we describe a new human thioredoxin peroxidase, antioxidant enzyme AOE372, identified by virtue of its protein-protein interaction with the product of a proliferation association gene, pag, which is also a thiol-specific

peroxidase, antioxidant enzyme AOE372, identified by virtue of its protein-protein interaction with the product of a **proliferation** association gene, pag, which is also a **thiol**-specific antioxidant. In human cells, AOE372 defines a **redox** pathway that specifically regulates NF-kappaB activity via a modulation of IkappaB-alpha phosphorylation in the cytoplasm. We show that AOE372 activity is regulated through either homo- or heterodimerization with other **thiol** peroxidases, implicating subunit assortment as a mechanism for regulating antioxidant specificities. AOE372 function suggests **thioredoxin** peroxidase as an immediate regulator of H2O2-mediated activation of NF-kappaB.

L38 ANSWER 40 OF 67 MEDLINE on STN DUPLICATE 18

ACCESSION NUMBER: 97400545 MEDLINE DOCUMENT NUMBER: PubMed ID: 9252380

TITLE: Generation of angiostatin by reduction and proteolysis of

plasmin. Catalysis by a plasmin reductase secreted by

cultured cells.

AUTHOR: Stathakis P; Fitzgerald M; Matthias L J; Chesterman C N;

Hogg P J

CORPORATE SOURCE: Centre for Thrombosis and Vascular Research, School of

Pathology and Department of Haematology, Prince of Wales Hospital, University of New South Wales, Sydney NSW 2052,

Australia.

SOURCE: Journal of biological chemistry, (1997 Aug 15) 272 (33)

20641-5.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH:

199709

ENTRY DATE:

Entered STN: 19970916

Last Updated on STN: 19970916 Entered Medline: 19970904

Extracellular manipulation of protein disulfide bonds has been implied in AB diverse biological processes, including penetration of viruses and endotoxin into cells and activation of certain cytokine receptors. demonstrate reduction of one or more disulfide bonds in the serine proteinase, plasmin, by a reductase secreted by Chinese hamster ovary or HT1080 cells. Reduction of plasmin disulfide bond(s) triggered proteolysis of the enzyme, generating fragments with the domain structure of the angiogenesis inhibitor, angiostatin. Two of the known reductases secreted by cultured cells are protein disulfide isomerase and thioredoxin, and incubation of plasmin with these purified reductases resulted in angiostatin fragments comparable with those generated from plasmin in cell culture. Thioredoxin-derived angiostatin inhibited proliferation of human dermal microvascular endothelial cells with half-maximal effect at approximately 0.2 microg/ml. Angiostatin made by cells and by purified reductases contained free sulfhydryl group(s), and S-carbamidomethylation of these thiol group(s) ablated biological activity. Neither protein disulfide isomerase nor thioredoxin were the reductases used by cultured cells, because immunodepletion of conditioned medium of these proteins did not affect angiostatin generating activity. The plasmin reductase secreted by HT1080 cells required a small cofactor for activity, and physiologically relevant concentrations of reduced qlutathione fulfilled this role. These results have consequences for plasmin activity and angiogenesis, particularly in the context of tumor growth and metastasis. Moreover, this is the first demonstration of extracellular reduction of a protein disulfide bond, which has general implications for cell biology.

L38 ANSWER 41 OF 67 MEDLINE on STN DUPLICATE 19

MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER:

PubMed ID: 9120263

TITLE:

Adult T cell leukemia (ATL)-derived factor/human thioredoxin prevents apoptosis of lymphoid cells induced by L-cystine and glutathione depletion:

possible involvement of thiol-mediated

redox regulation in apoptosis caused by pro-oxidant

state.

97240750

AUTHOR:

Iwata S; Hori T; Sato N; Hirota K; Sasada T; Mitsui A;

Hirakawa T; Yodoi J

CORPORATE SOURCE:

Department of Biological Responses, Kyoto University,

Japan.

SOURCE:

Journal of immunology (Baltimore, Md. : 1950), (1997 Apr 1)

158 (7) 3108-17.

Journal code: 2985117R. ISSN: 0022-1767.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: ENTRY MONTH:

Abridged Index Medicus Journals; Priority Journals

199704

ENTRY DATE:

Entered STN: 19970506

Last Updated on STN: 20000303 Entered Medline: 19970424

AB Thiol compounds, such as L-cysteine and glutathione (GSH), play crucial roles in the regulation of lymphocyte

proliferation. In this study, we analyzed the effect of L-cystine

and GSH depletion on lymphocyte survival and investigated the regulatory roles of adult T cell leukemia (ATL)-derived factor (ADF)/human thioredoxin (hTRX) in relation to these low m.w. thiols. MT-1, MT-2, and Jurkat cells underwent apoptosis when cultured in the L-cystine- and GSH-free medium within 18 to 24 h. Dichlorofluorescin oxidation assay indicated that the apoptosis in MT-1 and MT-2 cells was preceded by an increase in the level of intracellular hydrogen peroxide (H2O2). The addition of catalase and recombinant ADF/hTRX (rADF) partially blocked the apoptosis in a dose-dependent manner. rADF has been also shown to enhance the internalization of L-cystine into MT-2 cells in a dose-dependent manner, whereas oxidized rADF or mutated rADF that has no insulin-reducing activity failed to do so. Furthermore, culture in the L-cystine- and GSH-free medium lowered the cellular GSH content of PHA blasts, which was restored dose-dependently by rADF. These data suggest that the inability to neutralize oxidative stress results in the apoptosis of lymphoid cells under L-cystine- and GSH-depleted conditions. The protective effects of rADF may be explained by direct scavenging action on H2O2 (catalase-like activity) or by indirect neutralizing effects on the pro-oxidant status through enhancing the L-cystine internalization and elevating the intracellular GSH content.

L38 ANSWER 42 OF 67 MEDLINE ON STN ACCESSION NUMBER: 97160620 MEDLINE DOCUMENT NUMBER: PubMed ID: 9006954

TITLE: Characterization of a secretory type Theileria parva

glutaredoxin homologue identified by novel

screening procedure.

AUTHOR: Ebel T; Middleton J F; Frisch A; Lipp J

CORPORATE SOURCE: Vienna International Research Cooperation Center,

University of Vienna, A-1235 Vienna, Austria.

SOURCE: Journal of biological chemistry, (1997 Jan 31) 272 (5)

3042-8.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals OTHER SOURCE: GENBANK-U48417

ENTRY MONTH: 199703

ENTRY DATE: Entered STN: 19970321

Last Updated on STN: 19970321 Entered Medline: 19970313

ΑB The schizont stage of the protozoan parasite Theileria parva induces features characteristic of tumor cells in infected bovine T-cell lines. Most strikingly T. parva-infected cell lines acquire unlimited growth potential in vitro. Their proliferative state is entirely dependent on the presence of a viable parasite within the host cell cytoplasm. It has been postulated that parasite proteins either secreted into the host cell or expressed on the parasite surface membrane are involved in the parasite-host cell interaction. We used an in vitro transcription-translation-membrane translocation system to identify T. parva-derived cDNA clones encoding secretory or membrane proteins. Within 600 clones we found one encoding a 17-kDa protein which is processed by microsomal membranes to a 14-kDa protein (11E), presumably by signal peptidase. The processed form is expressed in the T-cell line TpM803 harboring viable parasites. By immunolocalization we show that the 11E protein mostly resides within the parasite, often in close vicinity to membranous structures, but in addition it appears at the surface membrane. Amino acid sequence comparison suggests that 11E belongs to the

glutaredoxin family, but is unique so far in containing a signal sequence for endoplasmic reticulum membrane translocation.

L38 ANSWER 43 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 1998048667 EMBASE

ACCESSION NOMBER. 1770040007 EMBASE

TITLE: Luminal peroxides in intestinal thiol-disulfide

balance and cell turnover.

AUTHOR: Tak Yee Aw

CORPORATE SOURCE: T.Y. Aw, Dept. of Molecular/Cellular Physiol, LSU Medical

Center, Shreveport, LA, United States

SOURCE: Comparative Biochemistry and Physiology - B Biochemistry

and Molecular Biology, (1997) Vol. 118, No. 3,

pp. 479-485. Refs: 65

ISSN: 0305-0491 CODEN: CBPBB8

PUBLISHER IDENT.: S 0305-0491(97)00220-4

COUNTRY: United States

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 002 Physiology

029 Clinical Biochemistry

048 Gastroenterology

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 19980227

Last Updated on STN: 19980227

AB Dietary intake of highly polyunsaturated fats represents a major source of lipid hydroperoxides in the intestinal lumen. Under conditions of high peroxide intake, excessive concentrations of lipid hydroperoxides can persist in the gut lumen and contribute to impairment of mucosal GSH-dependent detoxication pathways, enterocyte dysfunction independent of cell injury, and development of gut pathologies, including cancer.

This paper summarizes our current knowledge of the determinants of intestinal lipid hydroperoxide metabolism and of the physiological and biochemical processes in lipid peroxide-mediated changes in intestinal redox status, regulation of mucosal thiol and

antioxidant balance and control of intestinal cell turnover. This discussion is pertinent to understanding dietary peroxides and thiol redox balance in intestinal physiology and pathophysiology and the potential benefit of oral GSH in preserving

pathophysiology and the potential benefit of oral GSH in preserving metabolic integrity of the intestinal epithelium.

L38 ANSWER 44 OF 67 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED. on STN

ACCESSION NUMBER: 97137031 EMBASE

DOCUMENT NUMBER: 1997137031

TITLE: Redox state changes in density-dependent

regulation of proliferation.

AUTHOR: Hutter D.E.; Till B.G.; Greene J.J.

CORPORATE SOURCE: J.J. Greene, Department of Biology, Catholic University of

America, Washington, DC 20064, United States

SOURCE: Experimental Cell Research, (1997) Vol. 232, No.

2, pp. 435-438.

Refs: 25

ISSN: 0014-4827 CODEN: ECREAL

COUNTRY: United States
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 002 Physiology

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 970527

Last Updated on STN: 970527

The ability of certain transcription factors to bind to DNA has been ΔR demonstrated to be influenced by the redox environment. Therefore, fluctuations in the redox state of the cell may regulate the transcription of genes which control proliferation. To assess whether changes in the redox state may be related to proliferation, levels of oxidized (GSSG) and reduced (GSH) glutathione, the primary modulators of the redox state, were measured in cultures of varying densities of normal human fibroblasts which exhibit contact inhibition of proliferation, as well as fibrosarcoma cells, which lack this mechanism of growth control. Redox potentials calculated from normal, proliferating fibroblasts were found to be -34 mV more reducing than confluent, contact-inhibited cells. However, fibrosarcoma cells did not demonstrate this modulation in redox state. Further, to delineate whether these redox changes were the consequence or the cause of contact inhibition, cultures of subconfluent proliferating fibroblasts were treated with modulators of glutathione synthesis. Buthionine sulfoximine, an inhibitor of GSH synthesis, induced a less reducing redox state and decreased proliferation. In contrast, GSH synthesis precursors caused a more reduced redox state and increased proliferation. Collectively, these results suggest an interrelationship between

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on STN

ACCESSION NUMBER: 97259007 EMBASE

redox state and growth control.

DOCUMENT NUMBER: 1997259007

TITLE: Nitric oxide and superoxide induced p53 and Bax accumulation during mesangial cell apoptosis.

AUTHOR: Sandau K.; Pfeilschifter J.; Brune B.

CORPORATE SOURCE: Dr. B. Brune, University of Erlangen-Numberg, Faculty of

Medicine, Loschgestrasse 8, 91054 Erlangen, Germany

SOURCE: Kidney International, (1997) Vol. 52, No. 2, pp.

378-386. Refs: 51

ISSN: 0085-2538 CODEN: KDYIA5

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

028 Urology and Nephrology

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 970918

Last Updated on STN: 970918

During **proliferative** glomerulonephritis, the early phase of mesangiolysis is linked to increased nitric oxide (NO·) production. NO· as well as superoxide (O2-) are inflammatory mediators that are generated by mesangial cells (MC) after cytokine stimulation. Added individually, both radicals induce MC apoptosis. However, the co-existence of a defined NO·/O2- ratio is cross-protective. Apoptosis is characterized by specific features such as chromatin condensation, DNA strand breaks, and the occurrence of apoptotic regulating proteins. The **tumor** suppressor p53 and Bax (Bcl-2 associated protein x) are considered to be classical death promotors,

which accumulate after toxic insults. To study p53 and Bax protein accumulation in NO· and/or O2- induced apoptosis, we used the NO-donor S- nitrosoglutathione (GSNO) and the redox cycler 2,3-dimethoxy-1,4-naphtoquione (DMNQ). Both agonists initiated DNA fragmentation in a concentration dependent manner associated with transient p53 and Bax up-regulation. Co- generation of NO·/O2- resulted not only in reduced DNA fragmentation, but also in decreased Bax accumulation. Comparable to the NO·/O2-co- generation, cytokines failed to induce apoptosis. In contrast, cytokines in combination with pyrrolidine dithiocarbamate, which blocks endogenous superoxide dismutase, allowed p53 and Bax accumulation as well as DNA fragmentation. Our results demonstrate p53 and Bax as early components in NO· and O2- induced rat MC apoptosis and point to the NO·/O2- interaction as a naturally occurring cell defense mechanism.

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on STN

ACCESSION NUMBER: 97260943 EMBASE

DOCUMENT NUMBER: 1997260943

TITLE: Redox control as a target for anticancer

drug development.

AUTHOR: Kirkpatrick D.L.

CORPORATE SOURCE: D.L. Kirkpatrick, Department of Chemistry, University of

Rgina, Regina, Sask. S4S 0A2, Canada

SOURCE: Current Pharmaceutical Design, (1997) Vol. 3, No.

3, pp. 305-322.

Refs: 302

ISSN: 1381-6128 CODEN: CPDEFP

COUNTRY: Netherlands

DOCUMENT TYPE: Journal; General Review

FILE SEGMENT: 016 Cancer

029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 970911

Last Updated on STN: 970911

AB Cells maintain an intracellular environment that is reducing in the face of all oxidizing extracellular environment. Regulated alterations in the intracellular redox state (redox signalling) can modulate events such as DNA synthesis, enzyme activation, selective gene expression and regulation of the cell cycle. The primary consequence of intracellular redox signalling is a change in the oxidation state of cysteine residues of key proteins. This review will examine a number of the cellular redox systems which are in place to control the redox state, including such proteins as glutathione and glutathione reductase. thioredoxin and thioredoxin reductase, the highly cysteine rich, metalothioneins and the Ref-1' protein which plays a role in the activity of AP-1 and NF-kB. Signalling processes will be identified which are dependent on the redox state of controlling proteins and are potential targets for drug development and include transcription factors whose activation is a prerequisite for growth faster stimulated growth. The development of drugs which exploit the cellular redox state has grown dramatically over the last few years as the understanding of cellular redox has burgeoned. This review will attempt to present the current state of knowledge of agents in this category including those which exploit the hypoxic cellular environment,

those which participate through antioxidant pathways and the evolving area of interest involving agents which alter the signalling process through protein thioalkylation.

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on STN

ACCESSION NUMBER: 97243511 EMBASE

DOCUMENT NUMBER: 1997243511

TITLE: Chemical-induced changes in intracellular redox

state and in apoptosis.

AUTHOR: Jajte J.M.

CORPORATE SOURCE: Dr. J.M. Jajte, Department of Physical Hazards, Nofer

Institute of Occupational Med., P.O. Box 199, 90-950 Lodz,

Poland

SOURCE: International Journal of Occupational Medicine and

Environmental Health, (1997) Vol. 10, No. 2, pp.

203-212. Refs: 44

ISSN: 1232-1087 CODEN: IOMHEZ

COUNTRY: Poland

DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 005 General Pathology and Pathological Anatomy

052 Toxicology

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 970829

Last Updated on STN: 970829

AB Necrosis and apoptosis are two ways by which cells die. A major concept of apoptosis is that it is a controlled process. From this concept it follows that cells contain a molecule or molecules which under specific, regulated circumstances mediate cell death. Recent data confirm that oxygen free radicals can be mediators of apoptosis. Chemicals could induce apoptosis due to reactive oxygen species production and changes in the intracellular redox state. Therefore, a complete understanding of the processes involved in apoptosis, and mechanisms of its manipulation, could provide novel strategies to the control of xenobiotic toxicity and give an impetus to design new therapeutic interventions.

L38 ANSWER 48 OF 67 JICST-EPlus COPYRIGHT 2005 JST on STN

ACCESSION NUMBER: 970501427 JICST-EPlus

TITLE: Retinoid induces growth inhibition of adult T-cell leukemia

cells.

AUTHOR: MIYATAKE J; MAEDA Y

CORPORATE SOURCE: Kinki Univ. School of Medicine, Osaka, JPN

SOURCE: Acta Med Kinki Univ, (1997) vol. 22, no. 1, pp. 111-121.

Journal Code: S0990A (Fig. 9, Tbl. 1, Ref. 32)

ISSN: 0386-6092

PUB. COUNTRY: Japan

DOCUMENT TYPE: Journal; Article

LANGUAGE: English STATUS: New

AB The effects of retinoic acid (RA) on the cell growth and expression of interleukin-2 (IL-2) receptor (IL-2RA/p55, Tac, CD25) by the human T lymphotropic virus type I positive (HTLV-I(+)) T cell lines, HUT102 and ATL-2, were investigated. Incubation of these cells with RA resulted in marked growth inhibition and down-regulation of CD25 expression. Four clones of HUT102 cell lines were established by limiting dilution, and RA was shown to inhibit the growth and CD25 expression in three of these

clones, but in the fourth. However, RA did not induce growth inhibition of the HTLV-I-negative T cell lines, MOLT-4 and Jurkat, and of normal lymphocytes that had been stimulated wih phytohemagglutinin. We hydothesized that the sensivity to retinoids depends on an imbalance in intracellular redox potential. To examine the effect of exogenous thiol compounds for the growth inhibition of HTLV-I(+) T cell lines induced by RA, these cell lines were cultured with serveral thiol compounds (ATL-derived factor, thioredoxin, L-cystine and glutathione (GSH)), following the addition of RA in thiolfree medium. Unexpectedly, thiol compounds alone, when added after RA, did not restore the growth inhibition of HTLV-I(+) T cell lines induced by RA. However, when those cells were preincubated with thiol compounds for 24 hrs, no RA-induced growth inhibition was observed. These findings suggest that intracellular reductive environments induced by thiol compounds are associated with resistance to RA of HTLV-I(+) T cells, and that thiol compounds may play an important role in HTLV-I(+) T cell proliferation. (author abst.)

L38 ANSWER 49 OF 67 MEDLINE on STN DUPLICATE 20

ACCESSION NUMBER: 97112983 MEDLINE DOCUMENT NUMBER: PubMed ID: 8943236

TITLE: Induction of p21 mediated by reactive oxygen species formed

during the metabolism of aziridinylbenzoquinones by HCT116

cells.

AUTHOR: Qiu X; Forman H J; Schonthal A H; Cadenas E

CORPORATE SOURCE: Department of Molecular Pharmacology and Toxicology, School

of Pharmacy, University of Southern California, Los

Angeles, California 90033, USA.

CONTRACT NUMBER: 1RO1 ES05423 (NIEHS)

SOURCE: Journal of biological chemistry, (1996 Dec 13) 271 (50)

31915-21.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199701

ENTRY DATE: Entered STN: 19970219

Last Updated on STN: 19970219 Entered Medline: 19970117

AB Aziridinylbenzoquinones are a group of antitumor agents that elicit cytotoxicity by generating either alkylating intermediates or reactive oxygen species. The mechanism of toxicity may not always, however, involve profound damage of cellular constituents, but may involve a cytostatic effect through interference with the cell cycle. In this context, we have examined the induction of the cell cycle inhibitor p21 (WAF1, CIP1, or sdil), whose overexpression suppresses the growth of various tumor cells, in human tumor cells metabolizing 3,6-diaziridinyl-1,4-benzoquinone (DZQ) and its C2,C5-substituted derivatives: 2,5-bis-(carboethoxyamino) (AZQ) and 2, 5-bis-2(hydroxyethylamino) (BZQ). Both DZQ and AZQ were effectively activated by HCT116 human colonic carcinoma cells; the activation of the former involved largely a dicoumarol-sensitive activity, whereas that of the latter appeared to be accomplished primarily by one-electron transfer reductases. BZQ was not a substrate for the dicoumarol-sensitive enzyme in HCT116 cells. Cellular activation of the first two quinones was associated with formation of oxygen-centered radicals as detected by EPR in conjunction with the spin trap 5,5'-dimethyl-l-pyrroline-N-oxide. The

redox transitions of DZQ involved hydroxyl radical formation and were strongly inhibited by catalase, whereas those of AZQ showed a strong superoxide anion component sensitive to superoxide dismutase. These signals were suppressed by N-acetylcysteine with concomitant production of a thiyl radical adduct. This suggests an effective electron transfer between the thiol and free radicals formed during the activation of these quinones. DZQ and AZQ induced significantly the expression of p21 in HCT116 cells, but a 10-fold higher concentration of AZO was required to achieve the level of induction elicited by DZQ. BZQ had little effect on p21 expression. p21 induction at both mRNA and protein levels correlated with the inhibition of either cyclin-dependent kinase activity or cell proliferation. p21 induction elicited by the above quinones was inhibited by Nacetylcysteine, whereas the non-sulfur analog, N-acetylalanine, was without effect. Catalase and superoxide dismutase did not effect p21 induction by aziridinylbenzoquinones in HCT116 cells, thus suggesting that extracellular sources of oxygen radicals generated by plasma membrane reductases have no influence in the expression of this gene. Hydrogen peroxide, a product of quinone redox cycling, elicited an increase of p21 mRNA levels in HCT116 and K562 human chronic myelogenous leukemia cells. The latter lacks p53, one of the activators of p21 transcription, thus suggesting that p21 expression can be accomplished in a p53-independent manner in these cells. This study suggests that p21 induction is mediated by an increase in the cellular steady-state concentration of oxygen radicals and that the greater effectiveness in p21 induction by DZQ may be related to its efficient metabolism by NAD(P)H:quinone oxidoreductase activity in HCT116 cells.

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on STN

ACCESSION NUMBER: 96303120 EMBASE

DOCUMENT NUMBER: 1996303120

TITLE: Selenite and selenate inhibit human lymphocyte growth via

different mechanisms.

AUTHOR: Spyrou G.; Bjornstedt M.; Skog S.; Holmgren A. CORPORATE SOURCE: Department of Bioscience, Center for Biotechnology,

Karolinska Institutet, S-141 57 Huddinge, Sweden

SOURCE: Cancer Research, (1996) Vol. 56, No. 19, pp.

4407-4412.

ISSN: 0008-5472 CODEN: CNREA8

COUNTRY: United States
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 016 Cancer

029 Clinical Biochemistry 037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 961021

Last Updated on STN: 961021

AB Selenium compounds like selenite and selenate have strung inhibitory effects, particularly on mammalian tumor cell growth by unknown mechanisms. We found that the addition of sodium selenite and sodium selenate inhibited the growth of human 3B6 and BL41 lymphocytes. Selenite was more potent because 10 µM selenite produced a growth inhibitory effect similar to that of 250 µM selenate. The mechanism of action of selenite and selenate appears to be different. 3B6 and BL41 cells treated with selenite accumulated in the S-phase; however, selenate caused an accumulation of cells in G2. Selenite- mediated growth inhibition was irreversible, although the effects of selenate could be reversed.

Selenite, in contrast to selenate, is efficiently reduced by the thioredoxin system (thioredoxin, thioredoxin reductase, and NADPH). At concentrations required to observe a similar effect on cell growth, the activity of thioredoxin reductase, recently shown to be a selenoprotein, increased in selenite-treated cells and decreased in selenate-treated cells. Ribonucleotide reductase activity was inhibited in an in vitro assay by selenite and selenodiglutathione but not by selenate. These results show that selenite and selenate use different mechanisms to inhibit cell growth.

L38 ANSWER 51 OF 67 MEDLINE on STN DUPLICATE 21

ACCESSION NUMBER: 97139536 MEDLINE DOCUMENT NUMBER: PubMed ID: 8986137

TITLE: Oxidative inactivation of thioredoxin as a

cellular growth factor and protection by a Cys73-->Ser

mutation.

AUTHOR: Gasdaska J R; Kirkpatrick D L; Montfort W; Kuperus M; Hill

S R; Berggren M; Powis G

CORPORATE SOURCE: Arizona Cancer Center, University of Arizona Health

Services Center, Tucson 85724-5024, USA.

CONTRACT NUMBER: CA48725 (NCI)

SOURCE: Biochemical pharmacology, (1996 Dec 13) 52 (11) 1741-7.

Journal code: 0101032. ISSN: 0006-2952.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199701

ENTRY DATE: Entered STN: 19970219

Last Updated on STN: 19970219 Entered Medline: 19970123

AB Thioredoxin (Trx) is a widely distributed redox protein that regulates several intracellular redox-dependent processes and stimulates the proliferation of both normal and tumor cells. We have found that when stored in the absence of reducing agents, human recombinant Trx undergoes spontaneous oxidation, losing its ability to stimulate cell growth, but is still a substrate for NADPH-dependent reduction by human thioredoxin reductase. is a slower spontaneous conversion of Trx to a homodimer that is not a substrate for reduction by thioredoxin reductase and that does not stimulate cell proliferation. Both conversions can be induced by chemical oxidants and are reversible by treatment with the thiol reducing agent dithiothreitol. SDS-PAGE suggests that Trx undergoes oxidation to monomeric form(s) preceding dimer formation. We have recently shown by X-ray crystallography that Trx forms a dimer that is stabilized by an intermolecular Cys73-Cys73 disulfide bond. A Cys73-->Ser mutant Trx (C73S) was prepared to determine the role of Cys73 in oxidative stability and growth stimulation. C73S was as effective as Trx in stimulating cell growth and was a comparable substrate for thioredoxin reductase. C73S did not show spontaneous or oxidant-induced loss of activity and did not form a dimer. The results suggest that Trx can exist in monomeric forms, some of which are mediated by Cys73 that do not stimulate cell proliferation but can be reduced by thioredoxin reductase. Cys73 is also involved in formation of an enzymatically inactive homodimer, which occurs on long term storage or by chemical oxidation. Thus, although clearly involved in protein inactivation, Cys73 is not necessary for the growth stimulating activity of Trx.

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on STN

96311210 EMBASE ACCESSION NUMBER:

DOCUMENT NUMBER: 1996311210

Nitric oxide donors suppress erythropoietin production in TITLE:

vitro.

AUTHOR: Schobersberger W.; Hoffmann G.; Fandrey J.

Physiologisches Institut I, Universitat Bonn, Nussallee CORPORATE SOURCE:

11,D-53115 Bonn, Germany

Pflugers Archiv European Journal of Physiology, (SOURCE:

1996) Vol. 432, No. 6, pp. 980-985.

ISSN: 0031-6768 CODEN: PFLABK

COUNTRY: Germany

Journal; Article DOCUMENT TYPE: Physiology FILE SEGMENT: 002 025 Hematology

> 037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 961106

Last Updated on STN: 961106

Many inflammatory diseases are associated with a hypoproliferative AB anaemia. Patients with this anaemia often present with serum erythropoietin (EPO) concentrations that are too low for the degree of their anaemia. Proinflammatory cytokines, in addition to their inhibitory effects on proliferation of erythroid progenitors, could contribute to the pathogenesis of this anaemia by reducing EPO production. Because several cytokines stimulate nitric oxide (NO) synthase we propose that nitric oxide might mediate the suppression of EPO production during inflammation. In order to test this hypothesis we investigated the effects of NO donors on 24-h hypoxia-induced EPO production in the hepatocellular carcinoma cell line HepG2. Following application of the NO donors sodium nitroprusside (SNP), 3-morpholinosydnonimine (SIN-1), and S-nitroso-N-acetyl-D, L-penicillamine (SNAP), EPO production was dose-dependently reduced: compared to the untreated control EPO production was lowered by 89% with SNP (1000 μM), by 66% with SIN-1 (1000 $\mu M)$, and by 72% with SNAP (500 $\mu M)$. In contrast, 8-bromo-cGMP did not inhibit EPO formation. Since pyrogallol (300 μ M) and H2O2 (250 μM) showed a comparable suppression of EPO synthesis, we propose that NO might affect EPO production either by a similar direct influence on the cellular redox state or via increasing the cellular content of reactive oxygen species.

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on STN

ACCESSION NUMBER: 96019392 EMBASE

DOCUMENT NUMBER: 1996019392

TITLE: The organization of the human GSTP1-1 gene promoter and its

response to retinoic acid and cellular redox

status.

AUTHOR: Xia C.; Hu J.; Ketterer B.; Taylor J.B.

CORPORATE SOURCE: Department of Molecular Pathology, University College

London, Windeyer Building, Cleveland Street, London W1P 6DB,

United Kingdom

SOURCE: Biochemical Journal, (1996) Vol. 313, No. 1, pp.

155-161.

ISSN: 0264-6021 CODEN: BIJOAK

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article FILE SEGMENT: 022 Human Genetics

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 960130

Last Updated on STN: 960130

High levels of expression of GSTP1-1 are associated with cell proliferation, embryogenesis and malignancy. Given the role of glutathione S-transferase (GST) in detoxication, it is possible that GSTP1-1 evolved specifically to protect proliferating cells and share regulatory mechanisms with other cellular genes which are involved in cell division and tumorigenesis. We have previously shown that the expression of GSTP1 is suppressed by retinoic acid (RA) in the presence of the retinoic acid receptor (RAR) as a result of decreased transcription from its promoter. Through deletion analysis, we show here that the RA-RAR-dependent repression is mediated by the region -73 to +8. Further mutation analysis of this region indicates that the DNA sequence required for RA-RAR-dependent repression co-localizes with a consensus activator protein-1 (AP1) site essential for the promoter activity. The degree of repression correlates with the residual activity of the AP1 site. There are two adjacent G/C boxes. The one immediately downstream from the AP1 site is not essential for the promoter activity, but mutation of the second, further downstream, impairs the promoter. On the other hand, mutation of either of these two G/C boxes has little effect on RA-RAR suppression. We also show that the expression of GSTP1 is regulated by the redox status of the cell. Using the chloramphenicol acetyltransferase assay system, we have demonstrated that treatment with H2O2 induced transcription from the promoter and that this effect can be blocked by pre-incubation with Nacetylcysteine (NAG). It was shown that the induction by H2O2, is mediated by trans-acting factor NF-κB (nuclear factor κB), via a putative NF-kB site, 'GGGACCCTCC', located from -96 to -86. Co-transfection with an NF- κ B expression construct increased the promoter activity, an effect which could be blocked by co-transfection with an $I\kappa B$ (MAD-3) expression construct. Deletion of the NF-κB site abolished the effect of both H2O2, and co-transfection of NF- κ B. Interestingly, **NAC** is also an inducer for GSTP1. The effect of NAC was shown to be mediated largely by the AP1 site, since mutation of this site abolished the induction by NAG.

L38 ANSWER 54 OF 67 MEDLINE on STN DUPLICATE 22

ACCESSION NUMBER: 96291625 MEDLINE DOCUMENT NUMBER: PubMed ID: 8726363

TITLE: Reduction-oxidation (redox) state regulation of

extracellular matrix metalloproteinases and tissue inhibitors in cardiac normal and transformed fibroblast

cells.

AUTHOR: Tyagi S C; Kumar S; Borders S

CORPORATE SOURCE: Department of Medicine, Dalton Cardiovascular Research

Center, University of Missouri-Columbia 65212, USA.

CONTRACT NUMBER: GM-48595 (NIGMS)

SOURCE: Journal of cellular biochemistry, (1996 Apr) 61 (1) 139-51.

Journal code: 8205768. ISSN: 0730-2312.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199610

ENTRY DATE: Entered STN: 19961025

Last Updated on STN: 20021218

Entered Medline: 19961015 Latent matrix metalloproteinases (MMPs) in normal myocardium are activated AB in end-stage heart failure. In vitro oxidized glutathione (GSSG) activates myocardial MMPs which contains a cysteine residue. vivo GSSG induce the collagen lysis and cardiac dilatation. To assess whether thiol and non-thiol reducing agents have direct effect on the interstitial human heart fibroblast (HHF) proliferation and MMP expression, HHF and polyoma virus transformed fibroblast cells were cultured with or without the thiol-containing reduced (GSH) or oxidized (GSSG) glutathiones, pyrrolidine dithiocarbamate (PDTC) and N-acetylcysteine (NAC), and non-thiol ascorbic acid. After 100 micrograms/ml (approximately 0.3 mM) GSH or PDTC treatment the proliferative (synthetic) phenotype of transformed fibroblast cells was changed to quiescent (contractile) phenotype. Also, after GSH, PDTC, and ascorbic acid treatment the medium was then analyzed for MMP activity by zymography. The results indicate reduction in MMP expression in transformed fibroblast cells after GSH and PDTC treatments and no effect after ascorbic acid treatment. Based on reverse zymography, we observed the level of tissue inhibitor of metalloproteinase (TIMP) at a decreased level in transformed cells. The effect of the reducing agent at the gene transcription was measured by estimating mRNA (Northern blot

analysis) of MMP and of TIMP in the cells that were cultured in medium in the presence and absence of GSH. These results indicate that GSH induces MMP-2 and MMP-1 expression in normal HHF and that GSH reduces MMP-2 and MMP-1 in transformed fibroblast cells. After the treatment, the TIMP-2 level was repressed in normal HHF and TIMP-2 level increased in transformed fibroblast cells. These events are dependent on the nuclear transcription factor activity on the collagenase promoter in normal HHF cells. On the other hand, in polyoma transform fibroblast cells these events are not dependent on this collagenase promoter. These results suggest that oxidative environment induces normal HHF cell proliferation, and the reducing agent decreases normal HHF cell

proliferation by inducing MMP and repressing TIMP gene transcription. In transformed cells reducing agents inhibit MMP expression and increase TIMP levels, which suggests a role of antioxidants in preventing tumorigenesis.

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ACCESSION NUMBER: 96295578 EMBASE

DOCUMENT NUMBER: 1996295578

TITLE: Cytoprotective agents for anthracyclines.

AUTHOR: Dorr R.T.

CORPORATE SOURCE: Arizona Cancer Center, 1515 N Campbell Ave, Tucson, AZ

58724, United States

SOURCE: Seminars in Oncology, (1996) Vol. 23, No. 4

SUPPL. 8, pp. 23-34.

ISSN: 0093-7754 CODEN: SOLGAV

COUNTRY: United States

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: Cancer 016

Pharmacology 030

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 961021

Last Updated on STN: 961021

Anthracycline-induced cardiotoxicity is believed to be related to the AB generation of reactive oxygen species by at least two mechanisms: enzymatic reduction of the quinone with subsequent redox cycling and/or formation of an iron-anthracycline complex capable of intramolecular reduction and redox cycling. Both pathways may lead to the production of superoxide anions and highly reactive metabolites, such as hydroxyl radicals and hydrogen peroxide. As a result, membrane lipid peroxidation may ensue, producing damage in tissues like the heart, which have low antioxidant defenses (superoxide dismutase glutathione and especially, glutathione-peroxidase). Pharmacologic methods of interrupting this cycle have involved numerous antioxidants, such as the sulfhydryls N-acetylcysteine and cysteamine, and the lipophilic vitamin alpha tocopherol. Unfortunately, none of these compounds has been proven to be cardioprotective in patients receiving doxorubitin. In contrast, the water-soluble d-isomer of the iron chelator razoxane, dexrazoxane or ICRF-187, has been shown to reduce doxorubicin-induced cardiomyopathy. This has afforded greater cumulative doses of doxorubicin to be safely administered. The cytoprotective effect is apparently limited to the heart since there is no effect on antitumor efficacy and, unfortunately, no reduction in gastrointestinal toxicity, and with a slight increase in myelosuppression. More recent preclinical studies have also demonstrated cardioprotective activity for the aminothiol amifostine (WR-2721). In vitro, this agent has been shown to scavenge superoxide anions and hydroxyl radicals, the latter effect mediated by the active (dephosphorylated) metabolite, WR-1065. In tumor-bearing mice, amifostine reduces the lethality of high doses of doxorubicin without affecting antitumor activity. Finally, in vitro studies in neonatal rat heart cells have shown direct evidence of anthracycline cardioprotection for both amifostine and WR-1065. Cytoprotective drug levels of either agent were limited to 2.0 $\mu g/mL$, which is one tenth of the achievable peak plasma levels in humans. At this concentration, a 15-minute sulfhydryl pretreatment significantly prevented doxorubicin-induced depressions of myocyte adenosine triphosphate levels. Overall, these studies suggest that amifostine may have cytoprotective activity against doxorubicin-induced cardiac damage. Animal studies in a chronically dosed doxorubicin model are indicated; if positive, clinical trials testing this hypothesis will be warranted.

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ACCESSION NUMBER: 95077360 EMBASE

DOCUMENT NUMBER: 1995077360

TITLE: Superoxide and hydrogen peroxide in relation to mammalian

cell proliferation.

AUTHOR: Burdon R.H.

CORPORATE SOURCE: Department Bioscience/Biotechnology, University of

Strathclyde, Glasgow G4 ONR, United Kingdom

SOURCE: Free Radical Biology and Medicine, (1995) Vol.

18, No. 4, pp. 775-794.

ISSN: 0891-5849 CODEN: FRBMEH

COUNTRY: United States

DOCUMENT TYPE: Journal; General Review

FILE SEGMENT: 029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 950405

Last Updated on STN: 950405

AB A wide variety of normal and malignant cell types generate and release

superoxide or hydrogen peroxide in vitro either in response to specific cytokine/growth factor stimulus or constitutively in the case of tumour cells. These species at submicromolar levels appear to act as novel intra and intercellular 'messengers' capable of promoting growth responses in culture. The mechanisms may involve direct interaction with specific receptors or oxidation of growth signal transduction molecules such as protein kinases, protein phosphatases, transcription factors, or transcription factor inhibitors. It is also possible that hydrogen peroxide may modulate the redox state and activity of these important signal transduction proteins indirectly through changes in cellular levels of GSH and GSSG. Critical balances appear to exist in relation to cell proliferation on one hand and lipid peroxidation and cell death on the other. Progression to a more prooxidant state whilst initially leading to enhanced proliferative responses results subsequently in increased cell death.

L38 ANSWER 57 OF 67 MEDLINE on STN DUPLICATE 23

ACCESSION NUMBER: 95136249 MEDLINE DOCUMENT NUMBER: PubMed ID: 7834639

TITLE: High rates of thioredoxin secretion correlate

with growth arrest in hepatoma cells.

AUTHOR: Rubartelli A; Bonifaci N; Sitia R

CORPORATE SOURCE: Laboratory of Clinical Pathology, National Institute for

Cancer Research, Genova, Italy.

SOURCE: Cancer research, (1995 Feb 1) 55 (3) 675-80.

Journal code: 2984705R. ISSN: 0008-5472.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199502

ENTRY DATE: Entered STN: 19950314

Last Updated on STN: 19970203 Entered Medline: 19950228

AB Thioredoxin (TRX), a disulfide-reducing intracellular dithiol enzyme, is synthesized by both normal liver cells and the hepatocarcinoma cell line HepG2. Only the former, however, secrete abundant TRX extracellularly. When cultured in mild reducing conditions, HepG2 cells but not normal hepatocytes increase the rate of TRX secretion and undergo growth inhibition accompanied by morphological changes. Also, recombinant TRX inhibits proliferation of HepG2 cells. In contrast, exogenous thiols and TRX stimulate proliferation of a B-cell lymphoma line, indicating that different cell types respond differently to variations in the extracellular redox potential.

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on STN

ACCESSION NUMBER: 95189904 EMBASE

DOCUMENT NUMBER: 1995189904

TITLE: Increased levels of oxidized glutathione in CD4+

lymphocytes associated with disturbed intracellular redox balance in human immunodeficiency virus type

1 infection.

AUTHOR: Aukrust P.; Svardal A.M.; Muller F.; Lunden B.; Berge R.K.;

Ueland P.M.; Froland S.S.

CORPORATE SOURCE: Clinical Immunol./Infect. Dis. Sec., Medical Department A,

Rikshospitalet, N-0027 Oslo, Norway

SOURCE: Blood, (1995) Vol. 86, No. 1, pp. 258-267.

ISSN: 0006-4971 CODEN: BLOOAW

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 026 Immunology, Serology and Transplantation

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 950712

Last Updated on STN: 950712

We investigated the intracellular glutathione redox AB status in isolated lymphocyte subpopulations and monocytes in patients with human immunodeficiency virus type 1 (HIV-1) infection and in healthy controls. CD4+ lymphocytes from HIV-1-infected patients were primarily characterized by a substantial increase in oxidized glutathione levels and a considerable decrease in the ratio of reduced to total glutathione, in most cases below 0.5 in patients with symptomatic HIV-1 infection, rather than decreased levels of reduced glutathione. The increase in oxidized glutathione was strongly correlated with low numbers of CD4+ lymphocytes in peripheral blood and impaired stimulated interleukin-2 production and proliferation in peripheral blood mononuclear cells, which is compatible with an immunopathogenic role for these redox disturbances. The HIV-1-infected patients with the most advanced clinical and immunologic disease were also characterized by an increase in levels of reduced glutathione in monocytes, suggesting that the glutathione redox cycle may be differentially regulated in CD4+ lymphocytes and monocytes. We could not confirm previous reports suggesting cysteine deficiency as a major cause of disturbed

suggesting cysteine deficiency as a major cause of disturbed glutathione homeostasis during HIV-1 infection. The demonstrated glutathione abnormalities were correlated with raised serum levels of tumor necrosis factor α . These findings suggest that a therapeutical approach, which can restore the glutathione

redox dysbalance in CD4+ lymphocytes and decrease the inflammatory stress, may be worthwhile exploring in HIV-1 infection.

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on STN

ACCESSION NUMBER: 95081084 EMBASE

DOCUMENT NUMBER: 1995081084

TITLE: Analysis of studies related to tumorigenicity

induced by hydroguinone.

AUTHOR: Whysner J.; Verna L.; English J.C.; Williams G.M. CORPORATE SOURCE: Environmental Health/Safety Program, Division of

Pathology/Toxicology, American Health Foundation, Valhalla,

NY 10595, United States

SOURCE: Regulatory Toxicology and Pharmacology, (1995)

Vol. 21, No. 1, pp. 158-176. ISSN: 0273-2300 CODEN: RTOPDW

COUNTRY: United States

DOCUMENT TYPE: Journal; General Review

FILE SEGMENT: 016 Cancer

022 Human Genetics 025 Hematology

028 Urology and Nephrology

Occupational Health and Industrial Medicine Environmental Health and Pollution Control

048 Gastroenterology

052 Toxicology

LANGUAGE: English

SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 950412

Last Updated on STN: 950412

Hydroquinone (HQ) produced renal adenomas in male F344 rats, and these AB tumors appeared to arise from areas of spontaneous progressive nephropathy; the nephropathy itself has been found to be enhanced by HQ. Other neoplasms were not confirmed to be causally related to HQ among the reported bioassays. In the male F344 rat, HQ administered alone was not DNA reactive. HQ produced enhanced proliferation of renal tubular epithelium, presumably through toxicity involving glutathione conjugate formation. In the kidney, bone marrow, and other tissues, HQ may induce toxicity by redox cycling and lipid peroxidation. In bone marrow, HQ may produce microtubulin dysfunction, which is a plausible explanation for positive cytogenetic tests, the only consistently positive genotoxicity effect reported for HQ. Although HQ is a metabolic product of benzene, several lines of evidence suggest that the effects of HQ exposure are significantly different from those of benzene. Based upon the plausible mechanisms by which HQ may produce kidney tumors in male rats, we have concluded that occupational exposure levels of HQ are not predicted to be a cancer risk for humans.

L38 ANSWER 60 OF 67 MEDLINE on STN DUPLICATE 24

ACCESSION NUMBER: 94267168 MEDLINE DOCUMENT NUMBER: PubMed ID: 8207197

TITLE: Thiol-mediated redox regulation of

lymphocyte proliferation. Possible involvement of

adult T cell leukemia-derived factor and

glutathione in transferrin receptor expression.

AUTHOR: Iwata S; Hori T; Sato N; Ueda-Taniguchi Y; Yamabe T;

Nakamura H; Masutani H; Yodoi J

CORPORATE SOURCE: Department of Biological Responses, Kyoto University,

Japan.

SOURCE: Journal of immunology (Baltimore, Md.: 1950), (1994 Jun

15) 152 (12) 5633-42.

Journal code: 2985117R. ISSN: 0022-1767.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 199407

ENTRY DATE: Entered STN: 19940721

Last Updated on STN: 20000303 Entered Medline: 19940713

The proliferative response of PBMC to PHA, Con A, OKT3 mAb and AB IL-2-dependent proliferation of PHA-blasts was examined in a thiol-free environment (cultured in a L-cystine- and GSH-free medium). [3H] TdR incorporation assay and cell cycle analysis revealed that stimulated PBMC could not enter the S phase when deprived of these thiol compounds. In thiol-free cultures, an increase in intracellular free Ca2+ concentration and IL-2R alpha-chain/p 55 (Tac) induction was still observed, whereas transferrin receptor induction was markedly reduced, suggesting that the proliferative response of mitogenically stimulated PBMC was arrested in the late G1 phase in which transferrin receptor is induced. In GSH-depleted cultures, a similar reduction of the proliferative response of PBMC and PHA-blasts was observed when the concentration of L-cystine was lowered, in a dose-dependent manner. Each reduction or loss of proliferative response was partially restored by supplementation of 2-ME or adult T cell leukemia-derived factor (ADF)/human thioredoxin which is

considered to be an endogenous dithiol-reducing factor. L-Cystine transport analysis showed that mitogenically stimulated PBMC and PHA blasts incorporated L-cystine, whereas resting PBMC did not. Furthermore, ADF as well as 2-ME exhibited an enhancing activity on the L-cystine transport in PHA blasts. Together with the fact that L-cystine transport is a limiting step in glutathione synthesis, these findings suggest that GSH and ADF might cooperate in the thiol -mediated redox regulation process and might also play key roles in cell cycle (late G1 to S) progression of activated lymphocytes.

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on STN

ACCESSION NUMBER: 94142819 EMBASE

DOCUMENT NUMBER: 1994142819

TITLE: Redox regulation of signal transduction: Tyrosine

phosphorylation and calcium influx.

AUTHOR: Staal F.J.T.; Anderson M.T.; Staal G.E.J.; Herzenberg L.A.;

Gitler C.; Herzenberg L.A.

CORPORATE SOURCE: Department of Genetics, Stanford Univ. School of

Medicine, Stanford, CA 94305, United States

SOURCE: Proceedings of the National Academy of Sciences of the

United States of America, (1994) Vol. 91, No. 9,

pp. 3619-3622.

ISSN: 0027-8424 CODEN: PNASA6

COUNTRY: United States
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 026 Immunology, Serology and Transplantation

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 940602

Last Updated on STN: 940602

AB Studies presented here show that altering the intracellular redox balance by decreasing glutathione levels profoundly affects early signal transduction events in human T cells. In a T-cell receptor (TCR) signaling model, short- term pretreatment with buthionine sulfoximine, which specifically decreases intracellular glutathione, essentially abrogates the stimulation of calcium influx by anti-CD3 antibodies without significantly impairing other aspects of TCR-initiated signal transduction, such as overall levels of TCR- stimulated tyrosine phosphorylation. In an inflammatory-cytokine signaling model, the failure of tumor necrosis factor α to stimulate more than minimal tyrosine phosphorylation in lymphocytes is overcome by buthionine sulfoximine pretreatment-i.e., tumor necrosis factor α stimulates extensive tyrosine phosphorylation in glutathione-depleted lymphocytes. These redox-dependent changes in T-cell responsiveness suggest that the glutathione deficiency that we and others have demonstrated in human immunodeficiency virus-infected individuals may contribute significantly to the immunodeficiency and the increased inflammatory reactions in these individuals.

L38 ANSWER 62 OF 67 MEDLINE on STN DUPLICATE 25

ACCESSION NUMBER: 95019213 MEDLINE DOCUMENT NUMBER: PubMed ID: 7933622

TITLE: Measurement of ADF/thioredoxin in human serum and

its clinical significance.

AUTHOR: Kitaoka Y; Sachi Y; Mori T; Yodoi J

CORPORATE SOURCE: Department of Biological Responses, Kyoto University.

SOURCE: Rinsho byori. Japanese journal of clinical pathology, (1994

Aug) 42 (8) 853-9.

Journal code: 2984781R. ISSN: 0047-1860.

PUB. COUNTRY: Japan

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: Japanese

FILE SEGMENT: Priority Journals; AIDS

ENTRY MONTH: 199411

ENTRY DATE: Entered STN: 19941222

Last Updated on STN: 19941222 Entered Medline: 19941104

The adult T cell leukemia (ATL)-derived factor (ADF) was first described AB as an interleukin 2 receptor alpha chain (IL-2R alpha) inducing factor which is produced by an HTLV-I infected cell line. Subsequent purification and gene cloning proved that it is a human homologue of a bacterial reducing coenzyme, thioredoxin (TRX). ADF/human TRX (hTRX) has multiple functions both in the extracellular and intracellular compartments, such as cytokine activity, dithiol-reducing activity and radical scavenging activity. ADF/hTRX can facilitate the interactions between the transcriptional factors and its target DNA sequences, which may result in the overexpression of IL -2R alpha in HTLV-I infected cells. Recently, we have detected the presence of ADF/hTRX in human serum (sADF) obtained from healthy volunteers using the insulin reducing assay and Western blotting analysis. Another endogenous redox regulator, glutathione (GSH) system, has long been studied for its relation to cell proliferation and activation. Our recent data showed that thiol compounds such as L-cysteine and GSH may be involved in the activation and cell cycle progression of stimulated lymphocytes. We have also found that ADF/hTRX promotes L-cysteine transport into the cells and increases intracellular GSH content, indicating the close association between ADF/hTRX and GSH systems. Redox regulation by ADF/hTRX and GSH systems seems to play an important role in regulating cell proliferation and activation. To assess the possible alteration of the sADF level in pathological conditions, such as viral infections, an ELISA system for ADF/hTRX was recently established using two different monoclonal antibodies against rADF. (ABSTRACT TRUNCATED AT 250 WORDS)

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on STN

ACCESSION NUMBER: 94220957 EMBASE

DOCUMENT NUMBER: 1994220957

TITLE: Redox reagents and staurosporine inhibit

stimulation of the transcription regulator NF- κB following tumour necrosis factor treatment of

chronic B-leukaemia cells.

AUTHOR: Jabbar S.A.B.; Hoffbrand A.V.; Wickremasinghe R.G.

CORPORATE SOURCE: Department of Haematology, Royal Free Hospital School

Medicine, Pond Street, London NW3 2QG, United Kingdom

SOURCE: Leukemia Research, (1994) Vol. 18, No. 7, pp.

523-530.

ISSN: 0145-2126 CODEN: LEREDD

COUNTRY: United Kingdom
DOCUMENT TYPE: Journal; Article
FILE SEGMENT: 016 Cancer
025 Hematology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

ENTRY DATE: Entered STN: 940803

Last Updated on STN: 940803

B-chronic lymphocytic leukaemia (B-CLL) and hairy cell leukaemia cells AB (HCL) are refractory to stimulation by several cytokines which activate normal B-cells. However, tumour necrosis factor (TNF) promotes the proliferation of these cells. TNF regulates some of its cellular responses via the transcription factor NF-kB. Using an electrophoretic mobility shift assay, we demonstrate that TNF treatment of B-CLL and HCL cells in vitro resulted in the augmentation of NF- κB levels. In haemopoietic cell lines, TNF induction of NF- κ B is mediated via the generation of reactive oxygen intermediates and by the activation of protein kinase C (PKC). We have used activators and inhibitors of these pathways to unravel TNF signalling in the cells of ten patients with B-CLL and two with HCL, using the increase in NF-κB levels following TNF treatment as an end point. Raising glutathione levels with N-acetyl cysteine substantially reduced $NF-\kappa B$ induction by TNF in two of four samples, as did treatment of cells with the antioxidant butylated-hydroxytoluene in all three samples tested. These data suggest that redox mechanisms are involved in TNF signalling in these cells. Treatment with the PKC activator phorbol myristate acetate failed to activate NF-kB suggesting that this enzyme does not mediate the induction of NF- κB in these cells. However, the protein kinase inhibitor staurosporine inhibited TNF induction of NF-kB in four of five samples, suggesting that staurosporine-sensitive protein kinases (other than PKC) are involved in the signalling pathway. Our results suggest that PKC-independent pathways, including pathways sensitive to redox reagents, mediate the induction of NF-κB by TNF in chronic B-leukaemia cells. Additionally, these data suggest that defects in PKC-mediated pathways may contribute to the general reluctance of B-CLL and HCL cells to respond to mitogenic signals.

L38 ANSWER 64 OF 67 MEDLINE on STN DUPLICATE 26

ACCESSION NUMBER: 94132729 MEDLINE PubMed ID: 7507967

DOCUMENT NUMBER:

TITLE: Inhibition of activation-induced death in T cell hybridomas

by thiol antioxidants: oxidative stress as a

mediator of apoptosis.

AUTHOR: Sandstrom P A; Mannie M D; Buttke T M

CORPORATE SOURCE: Department of Microbiology and Immunology, East Carolina

University, School of Medicine, Greenville, NC 27858.

SOURCE: Journal of leukocyte biology, (1994 Feb) 55 (2) 221-6. Journal code: 8405628. ISSN: 0741-5400.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199403

ENTRY DATE: Entered STN: 19940318

> Last Updated on STN: 19970203 Entered Medline: 19940308

AB N-Acetylcysteine (NAC) is a well established thiol antioxidant which, after uptake, deacylation and conversion to glutathione functions as both a redox buffer and a reactive oxygen intermediate scavenger. We report here that NAC completely blocks activation induced death and associated DNA fragmentation of myelin basic protein (MBP) specific T cell hybridomas. Conversely, NAC had very little effect on the antigen driven proliferation of a MBP specific T cell line similar to that from

which the hybridomas were derived. NAC displayed an analogous absolute inhibition of mitogen mediated activation induced death, even if added up to 3 h post activation. Although glutathione was as efficient as NAC at blocking activation induced death, dithiothreitol displayed minimal inhibition while L-cysteine had no effect at all. The observation that certain thiol antioxidants such as NAC and glutathione can completely block the activation induced death of T cell hybridomas implicates redox regulation in this process.

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on STN

ACCESSION NUMBER: 94169017 EMBASE

DOCUMENT NUMBER: 19

1994169017

TITLE:

Oxidative damage and repair in the developing nervous

system.

AUTHOR:

Verity M.A.

CORPORATE SOURCE:

Division of Neuropathology, Brain Research Institute, UCLA

Center for the Health Sciences, Los Angeles, CA 90024-1732,

United States

SOURCE:

NeuroToxicology, (1994) Vol. 15, No. 1, pp.

81-92.

008

ISSN: 0161-813X CODEN: NRTXDN

COUNTRY:

United States

DOCUMENT TYPE:

Journal; Conference Article

FILE SEGMENT:

Neurology and Neurosurgery

LANGUAGE: SUMMARY LANGUAGE: English English

ENTRY DATE:

Entered STN: 940629

Last Updated on STN: 940629

AB Excessive production of reactive oxygen species (ROS) is a recognized cause of cell injury. In contrast to such well recognized cell injury, oxidative stress plays a role in cell proliferation, differentiation and tumor promotion. This review examines the role of oxidative stress in initiating and promoting the establishment of normal or abnormal neuronal patterns and subsequent neurogenesis within the central and peripheral nervous system. In particular, the role of apoptosis in both normal and abnormal neuronal development and maturation will be examined with especial reference to the induction of apoptotic cell death following abusive ligand- induced ion movements. The interaction of oxidant stress and immediate-early response gene activation is discussed with further reference to the induction of apoptosis. While glutamate receptor activation appears mandatory for coordinate maturation and neuritogenesis, such neuronal survival and differentiation is intimately dependent upon the intracellular glutathione redox potential, maintained by cystine uptake. Selected examples of reactive oxygen species induced injury pertaining to developmental neurotoxicology are presented and include starvation, irradiation injury and glutamate excitotoxicity.

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on STN

ACCESSION NUMBER: 91203269 EMBASE

DOCUMENT NUMBER:

1991203269

TITLE:

Augmented anti-proliferative effect in combined

use of human lymphotoxin with a nitrosourea derivative,

ACNU, and the involvement of glutathione

redox cycle.

AUTHOR:

Mashiba H.; Matsunaga K.; Kakutani T.

Division of Immunology, National Kyushu Cancer Center, CORPORATE SOURCE:

3-1-1 Notame, Minami-ku, Fukuoka 815, Japan

International Journal of Immunopharmacology, (1991 SOURCE:

>) Vol. 13, No. 4, pp. 333-338. ISSN: 0192-0561 CODEN: IJIMDS

COUNTRY: United Kingdom DOCUMENT TYPE: Journal: Article

Immunology, Serology and Transplantation FILE SEGMENT: 026

> Pharmacology 030

Drug Literature Index 037

LANGUAGE: English SUMMARY LANGUAGE: English

Entered STN: 911216 ENTRY DATE:

Last Updated on STN: 911216

The cytotoxic or cytostatic effect of the combined use of human lymphotoxin (LT) with 1-(4-amino-2-methyl-5-pyrimidinyl)methyl-3-(2chloroethyl)-3-nitrosour ea hydrochloride (ACNU) on cells or Meth A tumor cells was studied. Simultaneous addition of LT derived from a human lymphoid cell line with ACNU (200 or 500 µg/ml) significantly augmented the cytotoxic effect. Similar augmented inhibition was obtained when LT was added to ACNU-treated L cells. The pre-treatment of Meth A tumor cells with ACNU (25 or 50 µg/ml) augmented recombinant human LT-mediated cytostasis. However, the addition of glutathione (1.0 mg/ml) to ACNU-treated Meth A tumor cells significantly nullified the augmented antiproliferative effect of LT (10 U/ml). These results suggest that augmentation of the anti-proliferative effect on tumor cells could be induced through the combined use of LT with ACNU by lowering the intracellular level of glutathione.

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on STN

ACCESSION NUMBER: 88144244 EMBASE

DOCUMENT NUMBER:

1988144244

TITLE:

Toxic effects of acute glutathione depletion by

buthionine sulfoximine and

dimethylfumarate on murine mammary carcinoma

cells.

AUTHOR:

Dethlefsen L.A.; Lehman C.M.; Biaglow J.E.; Peck V.M.

CORPORATE SOURCE: Department of Radiology, Section of Experimental Oncology,

University of Utah Health Sciences Center, Salt Lake City,

UT 84132, United States

SOURCE:

Radiation Research, (1988) Vol. 114, No. 2, pp.

215-224.

ISSN: 0033-7587 CODEN: RAREAE

COUNTRY:

United States

DOCUMENT TYPE:

Journal

FILE SEGMENT:

016 Cancer

023 Nuclear Medicine 030 Pharmacology

037 Drug Literature Index

LANGUAGE:

English

SUMMARY LANGUAGE:

English

ENTRY DATE:

Entered STN: 911211

Last Updated on STN: 911211

Glutathione (GSH) depletion to .simeq. 5% of control for 48 h or AR longer by 0.05 mM L-buthionine sulfoximine (BSO) led to appreciable toxicity for the 66 murine mammary carcinoma

cells growing in vitro [L.A. Dethlefsen et al., Int. J. Radiat. Oncol.

Biol. Phys. 12, 1157-1160 (1986)]. Such toxicity in normal, proliferating cells in vivo would be undesirable. Thus the toxic effects after acute GSH depletion to .simeq. 5% of control by BSO plus dimethylfumarate (DMF) were evaluated in these same 66 cells to determine if this anti-proliferative effect could be minimized. Two hours of 0.025 mM $\bar{\text{DMF}}$ reduced GSH to 45% of control, while 6 h of 0.05 mM BSO reduced it to 16%. However, BSO (6 h) plus DMF (2 h) and BSO (24 h) plus DMF (2 h) reduced GSH to 4 and 2%, respectively. The incorporation (15-min pulses) of radioactive precursors into protein and RNA were unaffected by these treatment protocols. In contrast, cell growth was only modestly affected, but the incorporation of [3H]thymidine into DNA was reduced to 64% of control by the BSO (24 h) plus DMF (2 h) protocol even though it was unaffected by the BSO (6 h) plus DMF (2 h) treatment. The cellular plating efficiencies from both protocols were reduced to .simeq. 75% of control cells. However, the aerobic radiation response, as measured by cell survival, was not modified at doses of either 4.0 or 8.0 Gy. The growth rates of treated cultures, after drug removal, quickly returned to control rates and the resynthesis of GSH in cells from both protocols was also rapid. The GSH levels after either protocol were slightly above control by 12 h after drug removal, dramatically over control (.simeq. 200%) by 24 h, and back to normal by 48 h. Thus even a relatively short treatment with BSO and DMF resulting in a GSH depletion to 2-5% of control had a marked effect on DNA synthesis and plating efficiency and a modest effect on cellular growth. One cannot rule out a direct effect of the drugs, but presumably the antiproliferative effects are due to a depletion of nuclear GSH with the subsequent inhibition of the GSH/glutaredoxin-mediated conversion of ribonucleotides to deoxyribonucleotides. However, even after extended treatment, upon drug removal, GSH was rapidly resynthesized and cellular DNA synthesis and growth quickly resumed.

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L42 ANSWER 1 OF 8 USPATFULL on STN
ACCESSION NUMBER:
                              2004:184092 USPATFULL
TITLE:
                              Nucleic acid and corresponding protein entitled 98P4B6
                              useful in treatment and detection of cancer
INVENTOR(S):
                              Raitano, Arthur B., Los Angeles, CA, UNITED STATES
                              Ge, Wangmao, Culver City, CA, UNITED STATES
                              Jakobovits, Aya, Beverly Hills, CA, UNITED STATES
                              Challita-Eid, Pia M., Encino, CA, UNITED STATES
                              Faris, Mary, Los Angeles, CA, UNITED STATES
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A1 20030404 (10) APPLICATION INFO.: US 2003-407484

Continuation-in-part of Ser. No. US 1999-455486, filed RELATED APPLN. INFO.: on 6 Dec 1999, PENDING Continuation-in-part of Ser. No.

US 1999-323873, filed on 1 Jun 1999, GRANTED, Pat. No.

US 6329503

DATE NUMBER

US 1998-91183P 19980630 (60) US 1998-87520P 19980601 (60) PRIORITY INFORMATION:

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DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

Kate H. Murashige, Morrison & Foerster LLP, Suite 500, LEGAL REPRESENTATIVE:

3811 Valley Centre Drive, San Diego, CA, 92130

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 97 Drawing Page(s)

LINE COUNT: 22646

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A novel gene 098P4B6 (also designated STEAP-2) and its encoded protein, and variants thereof, are described wherein 98P4B6 exhibits tissue specific expression in normal adult tissue, and is aberrantly expressed in the cancers listed in Table I. Consequently, 98P4B6 provides a diagnostic, prognostic, prophylactic and/or therapeutic target for cancer. The 98P4B6 gene or fragment thereof, or its encoded protein, or variants thereof, or a fragment thereof, can be used to elicit a humoral or cellular immune response; antibodies or T cells reactive with 98P4B6 can be used in active or passive immunization.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 2 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2003:294798 USPATFULL

TITLE: Manipulating nitrosative stress to kill pathologic

microbes, pathologic helminths and pathologically

proliferating cells or to upregulate

nitrosative stress defenses

INVENTOR(S): Stamler, Jonathan S., Chapel Hill, NC, UNITED STATES

Griffith, Owen W., Milwaukee, WI, UNITED STATES

NUMBER KIND DATE -----

US 2003207815 A1 20031106 US 2003-417238 A1 20030417 (10) PATENT INFORMATION: APPLICATION INFO.:

Division of Ser. No. US 2001-13455, filed on 13 Dec RELATED APPLN. INFO.: 2001, GRANTED, Pat. No. US 6608110 Continuation of Ser.

No. US 2000-690989, filed on 18 Oct 2000, GRANTED, Pat. No. US 6359004 Division of Ser. No. US 1999-361167, filed on 27 Jul 1999, GRANTED, Pat. No. US 6180824 Division of Ser. No. US 1997-852490, filed on 7 May

1997, GRANTED, Pat. No. US 6057367

NUMBER DATE

US 1996-25819P 19960830 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Eric S. Spector, JONES, TULLAR & COOPER, P.C., Eads

Station, P.O. Box 2266, Arlington, VA, 22202

NUMBER OF CLAIMS: 69 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT: 3394

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Mammals are treated for infections or for conditions associated with pathologically proliferating mammalian cell growth (for example certain cancers, restenosis, benign prostatic hypertrophy) by administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathologically proliferating mammalian cells. Novel agents include α -alkyl-S-alkyl-homocysteine sulfoximines wherein the α -alkyl contains 2 to 8 carbon atoms, and the S-alkyl-contains 1 to 10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g., humans at risk for a stroke because of having had a transient ischemic attack, are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 3 OF 8 USPATFULL on STN

ACCESSION NUMBER:

2003:258639 USPATFULL

TITLE:

207 human secreted proteins

INVENTOR(S):

Ni, Jian, Germantown, MD, UNITED STATES Ebner, Reinhard, Gaithersburg, MD, UNITED STATES

LaFleur, David W., Washington, DC, UNITED STATES Moore, Paul A., Germantown, MD, UNITED STATES

Olsen, Henrik S., Gaithersburg, MD, UNITED STATES Rosen, Craig A., Laytonsville, MD, UNITED STATES

Ruben, Steven M., Olney, MD, UNITED STATES

Soppet, Daniel R., Centreville, VA, UNITED STATES

Young, Paul E., Gaithersburg, MD, UNITED STATES Shi, Yanggu, Gaithersburg, MD, UNITED STATES

Florence, Kimberly A., Rockville, MD, UNITED STATES

Wei, Ying-Fei, Berkeley, CA, UNITED STATES

Florence, Charles, Rockville, MD, UNITED STATES Hu, Jing-Shan, Mountain View, CA, UNITED STATES

Li, Yi, Sunnyvale, CA, UNITED STATES

Kyaw, Hla, Frederick, MD, UNITED STATES

Fischer, Carrie L., Burke, VA, UNITED STATES Ferrie, Ann M., Painted Post, NY, UNITED STATES

Fan, Ping, Potomac, MD, UNITED STATES

Feng, Ping, Gaithersburg, MD, UNITED STATES

Endress, Gregory A., Florence, MA, UNITED STATES Dillon, Patrick J., Carlsbad, CA, UNITED STATES

Carter, Kenneth C., North Potomac, MD, UNITED STATES

Brewer, Laurie A., St. Paul, MN, UNITED STATES

Yu, Guo-Liang, Berkeley, CA, UNITED STATES Zeng, Zhizhen, Lansdale, PA, UNITED STATES

Greene, John M., Gaithersburg, MD, UNITED STATES

NUMBER KIND DATE

RELATED APPLN. INFO.:

PATENT INFORMATION: US 2003181692 A1 20030925 APPLICATION INFO.: US 2001-933767 A1 20010822 (9)

Continuation-in-part of Ser. No. WO 2001-US5614, filed on 21 Feb 2001, PENDING Continuation-in-part of Ser. No. US 1998-205258, filed on 4 Dec 1998, PENDING

	NUMBER	DATE		
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US 1997-70923P 19971218 (60)
US 1998-92921P 19980715 (60)
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US 1998-92921P 19980730 (60)
US 1998-94657P 19980730 (60)
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DOCUMENT TYPE:

Utility APPLICATION

FILE SEGMENT:

LEGAL REPRESENTATIVE: HUMAN GENOME SCIENCES INC, 9410 KEY WEST AVENUE,

ROCKVILLE, MD, 20850

NUMBER OF CLAIMS: 23 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 10 Drawing Page(s)

LINE COUNT: 32746

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel human secreted proteins and isolated nucleic acids containing the coding regions of the genes encoding such proteins. Also provided are vectors, host cells, antibodies, and recombinant methods for producing human secreted proteins. The invention further relates to diagnostic and therapeutic methods useful for diagnosing and treating diseases, disorders, and/or conditions related to these novel human secreted proteins.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 4 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2003:141028 USPATFULL

TITLE: MANIPULATING NITROSATIVE STRESS TO KILL PATHOLOGIC

MICROBES, PATHOLOGIC HELMINTHS AND PATHOLOGICALLY

PROLIFERATING CELLS OR TO UPREGULATE

NITROSATIVE STRESS DEFENSES

Stamler, Jonathan S., Chapel Hill, NC, UNITED STATES INVENTOR(S):

Griffith, Owen W., Milwaukee, WI, UNITED STATES

NUMBER KIND

US 2003096870 A1 20030522 PATENT INFORMATION:

B2 20030819 US 6608110

US 2001-13455 A1 20011213 (10) APPLICATION INFO.:

Continuation of Ser. No. US 2000-690989, filed on 18 RELATED APPLN. INFO.: Oct 2000, GRANTED, Pat. No. US 6359004 Division of Ser. No. US 1999-361167, filed on 27 Jul 1999, GRANTED, Pat.

No. US 6180824 Division of Ser. No. US 1997-852490, filed on 7 May 1997, GRANTED, Pat. No. US 6057367

> DATE NUMBER

US 1996-25819P 19960830 (60) PRIORITY INFORMATION: <--

DOCUMENT TYPE: Utility APPLICATION FILE SEGMENT:

Eric S. Spector, JONES, TULLAR & COOPER, P.C., Eads LEGAL REPRESENTATIVE:

Station, P.O. Box 2266, Arlington, VA, 22202

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT: 3394

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Mammals are treated for infections or for conditions associated with pathologically proliferating mammalian cell growth (for

example, certain cancers, restenosis, benign prostatic

hypertrophy) by administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathologically proliferating mammalian cells. Novel agents include α -alkyl-S-alkyl-homocysteine sulfoximines wherein the α -alkyl contains 2 to 8 carbon atoms, and the S-alkyl- contains 1 to 10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g., humans at risk for a stroke because of having had a transient ischemic attack, are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of

L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 5 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2002:57833 USPATFULL

TITLE: Manipulating nitrosative stress to upregulate

nitrosative stress defenses

INVENTOR(S): Stamler, Jonathan S., Chapel Hill, NC, United States

Griffith, Owen W., Milwaukee, WI, United States

PATENT ASSIGNEE(S): Duke University, Durham, NC, United States (U.S.

corporation)

The Medical College of Wisconsin, Milwaukee, WI, United

States (U.S. corporation)

KIND NUMBER DATE -----US 6359004 B1 20020319 US 2000-690989 20001018 (9) PATENT INFORMATION: APPLICATION INFO.:

Continuation of Ser. No. US 1999-361167, filed on 27 RELATED APPLN. INFO.:

Jul 1999, now patented, Pat. No. US 6180824 Division of

Ser. No. US 1997-852490, filed on 7 May 1997, now
patented, Pat. No. US 6057367

NUMBER DATE

PRIORITY INFORMATION: US 1996-25819P 19960830 (60) <--

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Weddington, Kevin E.

NUMBER OF CLAIMS: 2 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 2 Drawing Figure(s); 2 Drawing Page(s)

LINE COUNT: 3105

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Mammals are treated for infections or for conditions associated with pathologically proliferating mammalian cell growth (for example, certain cancers, restenosis, benign prostatic hypertrophy) by administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathologically proliferating mammalian cells. Novel agents include α-alkyl-S-alkyl-homocysteine sulfoximines wherein the α -alkyl contains 2 to 8 carbon atoms, and the S-alkyl- contains 1 to 10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g., humans at risk for a stroke because of having had a transient ischemic attack, are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 6 OF 8 USPATFULL on STN

ACCESSION NUMBER: 2001:14679 USPATFULL

TITLE: Manipulating nitrosative stress to kill pathologic

microbes, pathologic helminths and pathologically,

proliferating cells or to upregulate

nitrosative stress defenses

INVENTOR(S): Stamler, Jonathan S., Chapel Hill, NC, United States

Griffith, Owen W., Milwaukee, WI, United States

PATENT ASSIGNEE(S): Duke University, Durham, NC, United States (U.S.

corporation)

The Medical College of Wisconsin, Milwaukee, WI, United

States (U.S. corporation)

RELATED APPLN. INFO.: Division of Ser. No. US 1997-852490, filed on 7 May

1997, now patented, Pat. No. US 6057367

NUMBER DATE

PRIORITY INFORMATION: US 1996-25819P 19960830 (60) <--

DOCUMENT TYPE: Utility

FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Weddington, Kevin E.

NUMBER OF CLAIMS:

15

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

2 Drawing Figure(s); 2 Drawing Page(s)

LINE COUNT:

3128

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Mammals are treated for infection or for conditions associated with AB pathologically proliferating mammalian cell growth (for example, certain cancers, restenosis, benign prostatic hypertrophy) by administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathologically proliferating mammalian cells. Novel agents include α -alkyl-S-alkyl-homocysteine sulfoximines wherein the α -alkyl contains 2 to 8 carbon atoms, and the S-alkyl- contains 1 to 10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g., humans at risk for a stroke because of having had a transient ischemic attack, are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 7 OF 8 USPATFULL on STN

ACCESSION NUMBER:

2000:54150 USPATFULL

TITLE:

Manipulating nitrosative stress to kill pathologic microbes, pathologic helminths and pathologically

proliferating cells or to upregulate

L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

nitrosative stress defenses

INVENTOR (S):

Stamler, Jonathan S., Chapel Hill, NC, United States Griffith, Owen W., Milwaukee, WI, United States

PATENT ASSIGNEE(S):

Duke University, Durham, NC, United States (U.S.

corporation)

The Medical College of Wisconsin Research Foundation, Inc., Milwaukee, WI, United States (U.S. corporation)

NUMBER	KIND	DATE	
US 6057367		20000502	

PATENT INFORMATION: APPLICATION INFO.:

US 1997-852490 19970507 (8)

NUMBER DATE ______

PRIORITY INFORMATION:

US 1996-25819P 19960830 (60)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER:

Weddington, Kevin E.

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

2 Drawing Figure(s); 2 Drawing Page(s) LINE COUNT: 3415

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Mammals are treated for infections or for conditions associated with pathologically proliferating mammalian cell growth (for

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example, certain cancers, restenosis, benign prostatic hypertrophy) by administration of a manipulator of nitrosative stress to selectively kill or reduce the growth of the microbes or helminths causing the infection or of host cells infected with the microbes or of the pathologically proliferating mammalian cells. Novel agents include α -alkyl-S-alkyl-homocysteine sulfoximines wherein the $\alpha\text{-alkyl}$ contains 2 to 8 carbon atoms, and the S-alkyl-contains 1 to 10 carbon atoms. In another invention herein, mammals in need of increased nitrosative stress defenses are treated, e.g., humans at risk for a stroke because of having had a transient ischemic attack, are treated. Treatments to increase nitrosative stress defenses include, for example, repeated administrations of low doses of manipulators of nitrosative stress so that the subject treated has increased tolerance to nitrosative stress. In still another invention, mammals are treated for protozoal infections by systemic administration of L-buthionine-S-sulfoximine and agent that increases nitrosative stress.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L42 ANSWER 8 OF 8 USPATFULL on STN

ACCESSION NUMBER: 1998:156914 USPATFULL

TITLE: Compounds and methods for the diagnosis, treatment and

prevention of diseases of cell death

INVENTOR(S): Brown, Robert, Needham, MA, United States

Horvitz, H. Robert, Cambridge, MA, United States

Rosen, Daniel R., Dedham, MA, United States

PATENT ASSIGNEE(S): The General Hospital Corporation, Boston, MA, United

States (U.S. corporation)

Massachusetts Institute of Technology, Cambridge, MA,

United States (U.S. corporation)

NUMBER KIND DATE -----US 5849290 19981215 US 1995-486953 19950607 (8) PATENT INFORMATION:

APPLICATION INFO.:

Division of Ser. No. US 1994-204052, filed on 28 Feb RELATED APPLN. INFO.:

1994 which is a continuation-in-part of Ser. No. US

1993-23980, filed on 26 Feb 1993

DOCUMENT TYPE: Utility Granted FILE SEGMENT: PRIMARY EXAMINER: Grimes, Eric LEGAL REPRESENTATIVE: Clark & Elbing LLP

NUMBER OF CLAIMS: 6 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 30 Drawing Figure(s); 17 Drawing Page(s)

LINE COUNT: 2365

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Disclosed is the family of genes responsible for the neurodegenerative diseases, particularly Amyotrophic Lateral Sclerosis. Methods and compounds for the diagnosis, prevention, and therapy of the disease are

also disclosed.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Spivack 09/913,435

=> d his ful

L37

(FILE 'HOME' ENTERED AT 16:52:53 ON 17 SEP 2005)

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               1 SEA ABB=ON PDTC/CN
L2
               0 SEA ABB=ON 2,3-DIMERCAPTO-1-PROPANE SULFONIC ACID/CN
L3
              1 SEA ABB=ON 2,3-DIMERCAPTO-1-PROPANESULFONIC ACID/CN
L4
              1 SEA ABB=ON DITHIOCARBAMATE/CN
L5
              1 SEA ABB=ON DITHIOTHREITOL/CN
L6
              O SEA ABB=ON GLUTATHIONE ESTER/CN
L7
              1 SEA ABB=ON BUTHIONINE SULFOXIMINE/CN
1 SEA ABB=ON METHIONINE SULFOXIMINE/CN
0 SEA ABB=ON N-ACETYL CYSTEINE/CN
L8
L9
L10
L11
              1 SEA ABB=ON N-ACETYLCYSTEINE/CN
             1 SEA ABB=ON CYSTEAMINE/CN
2 SEA ABB=ON LIPOIC ACID/CN
L12
L13
             1 SEA ABB=ON THIOCTIC ACID/CN
L14
             0 SEA ABB=ON 2-MERCAPTO-1-PROPANESULFONIC ACID/CN
L15
             3 SEA ABB=ON DMSA/CN
L16
             1 SEA ABB=ON 304-55-2/RN
L17
              1 SEA ABB=ON MESNA/CN
L18
L19
              1 SEA ABB=ON DITHIOTHREITOL/CN
L20
              O SEA ABB=ON ACIVACIN/CN
              1 SEA ABB=ON ACIVICIN/CN
L21
     FILE 'HCAPLUS' ENTERED AT 17:04:07 ON 17 SEP 2005
               1 SEA ABB=ON ACIVACIN
L22
     FILE 'REGISTRY' ENTERED AT 17:07:30 ON 17 SEP 2005
L23
              1 SEA ABB=ON ACIVICIN/CN
             17 SEA ABB=ON L1 OR L2 OR L4 OR L5 OR L6 OR L8 OR L9 OR L11 OR
L24
                 L12 OR L13 OR L14 OR L16 OR L17 OR L18 OR L19 OR L21 OR L23
     FILE 'HCAPLUS' ENTERED AT 17:08:34 ON 17 SEP 2005
          23859 SEA ABB=ON L24
L25
         146867 SEA ABB=ON L25 OR (?ETHACRYNIC? OR (2,3-DIMERCAPTO-1-PROPANESU
L26
                 LFONIC? OR 2-MERCAPTO-1-PROPANESULFONIC) (W) ?ACID? OR ?DITHIOCAR
                 BAMATE? OR ?DITHIOTHREITOL? OR ?GLUTATHIONE? OR (?BUTHIONINE?
                 OR ?METHIONINE?) (W) ?SULFOXIMINE? OR N-?ACETYLCYSTEINE? OR NAC
                 OR ?CYSTEAMINE?)
         149253 SEA ABB=ON L26 OR (?LIPOIC? OR ?THIOCTIC? OR 2-MERCAPTO-1-PROP
L27
                 ANESULFONIC?) (W) ?ACID? OR DMSA OR MESNA OR ?REDUC? (W) ?CYSTEINE?
                  OR ?ACIVACIN? OR ?ACIVICIN?
L28
           7586 SEA ABB=ON L27 AND ?REDOX?
           1110 SEA ABB=ON L28 AND (?CANCER? OR ?CARCIN? OR ?NEOPLAS? OR
L29
                 ?TUMOR? OR ?TUMOUR?)
L30
            199 SEA ABB=ON L29 AND ?PROLIFERAT?
             48 SEA ABB=ON L30 AND ?THIOL?
L31
              0 SEA ABB=ON L30 AND NON? (W) ?VIRAL?
L32
             57 SEA ABB=ON L30 AND (2VIRUS? OR 7VIRAL?)
54 SEA ABB=ON L30 AND (PRD<19990216 OR PD<19990216)
L33
L34
             85 SEA ABB=ON L31 OR L34
L35
             54 SEA ABB=ON L35 AND (PRD<19990216 OR PD<19990216)
L36
     FILE 'MEDLINE, CANCERLIT, EMBASE, JAPIO, JICST-EPLUS' ENTERED AT 17:24:57
     ON 17 SEP 2005
```

103 SEA ABB=ON L35

L38 67 DUP REMOV L37 (36 DUPLICATES REMOVED)

FILE 'USPATFULL' ENTERED AT 17:27:05 ON 17 SEP 2005

L39 808 SEA ABB=ON L35 AND (PRD<19990216 OR PD<19990216)

L40 316 SEA ABB=ON L39 AND ?THIOL?

L41 140 SEA ABB=ON L40 AND ?CHEMOTHERAPEUTIC?(W)?AGENT?

L42 8 SEA ABB=ON L41 AND NON? (W) ?VIRAL?

SAV L42 SPI435L42/A

FILE 'HCAPLUS' ENTERED AT 17:29:19 ON 17 SEP 2005 SAV L36 SPI435L36/A

FILE 'MEDLINE, CANCERLIT, EMBASE, JAPIO, JICST-EPLUS' ENTERED AT 17:29:39 ON 17 SEP 2005

SAV L38 SPI435L38/A

FILE HOME

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 16 SEP 2005 HIGHEST RN 863378-74-9 DICTIONARY FILE UPDATES: 16 SEP 2005 HIGHEST RN 863378-74-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

Experimental and calculated property data are now available. For more information enter HELP PROP at an arrow prompt in the file or refer to the file summary sheet on the web at: http://www.cas.org/ONLINE/DBSS/registryss.html

FILE HCAPLUS

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FILE COVERS 1907 - 17 Sep 2005 VOL 143 ISS 13 FILE LAST UPDATED: 16 Sep 2005 (20050916/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE MEDLINE

FILE LAST UPDATED: 16 SEP 2005 (20050916/UP). FILE COVERS 1950 TO DATE.

On December 19, 2004, the 2005 MeSH terms were loaded.

The MEDLINE reload for 2005 is now available. For details enter HELP RLOAD at an arrow promt (=>). See also:

http://www.nlm.nih.gov/mesh/

http://www.nlm.nih.gov/pubs/techbull/nd04/nd04_mesh.html

OLDMEDLINE now back to 1950.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2005 vocabulary.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE CANCERLIT

FILE COVERS 1963 TO 15 Nov 2002 (20021115/ED)

On July 28, 2002, CANCERLIT was reloaded. See HELP RLOAD for details.

CANCERLIT thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2002 vocabulary. Enter HELP THESAURUS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE EMBASE

FILE COVERS 1974 TO 15 Sep 2005 (20050915/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE JAPIO

FILE LAST UPDATED: 5 SEP 2005 <20050905/UP>
FILE COVERS APR 1973 TO APRIL 28, 2005

<<< GRAPHIC IMAGES AVAILABLE >>>

FILE JICST-EPLUS

FILE COVERS 1985 TO 13 SEP 2005 (20050913/ED)

THE JICST-EPLUS FILE HAS BEEN RELOADED TO REFLECT THE 1999 CONTROLLED TERM (/CT) THESAURUS RELOAD.

FILE USPATFULL
FILE COVERS 1971 TO PATENT PUBLICATION DATE: 15 Sep 2005 (20050915/PD)
FILE LAST UPDATED: 15 Sep 2005 (20050915/ED)
HIGHEST GRANTED PATENT NUMBER: US6944881
HIGHEST APPLICATION PUBLICATION NUMBER: US2005204445
CA INDEXING IS CURRENT THROUGH 15 Sep 2005 (20050915/UPCA)
ISSUE CLASS FIELDS (/INCL) CURRENT THROUGH: 15 Sep 2005 (20050915/PD)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Jun 2005
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2005

>>> USPAT2 is now available. USPATFULL contains full text of the <<< >>> original, i.e., the earliest published granted patents or <<< >>> applications. USPAT2 contains full text of the latest US >>> publications, starting in 2001, for the inventions covered in <<< <<< >>> USPATFULL. A USPATFULL record contains not only the original >>> published document but also a list of any subsequent <<< >>> publications. The publication number, patent kind code, and <<< >>> publication date for all the US publications for an invention <<< >>> are displayed in the PI (Patent Information) field of USPATFULL <<< >>> records and may be searched in standard search fields, e.g., /PN, <<< >>> /PK, etc. <<< >>> USPATFULL and USPAT2 can be accessed and searched together <<< >>> through the new cluster USPATALL. Type FILE USPATALL to <<< >>> enter this cluster. <<< >>> <<< >>> Use USPATALL when searching terms such as patent assignees, <<< >>> classifications, or claims, that may potentially change from <<< >>> the earliest to the latest publication. <<<

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 14 September 2005 (20050914/ED)

FILE RELOADED: 19 October 2003.